

HANDBOOK OF DIGESTIVE DISEASES

HANDBOOK

OF

DIGESTIVE DISEASES

BY

JOHN L. KANTOR M.D., F.A.C.P.

Late Associate in Medicine Columbia University Gastroenterologist and
Associate Roentgenologist Montefiore Hospital,
New York

AND

ANTHONY M. KASICH, M.D., F.A.C.P.

Lecturer in Medicine Columbia University Adjunct Physician
Montefiore Hospital Assistant Visiting Physician Bellevue
Hospital Assistant Adjunct Gastroenterologist
Lenox Hill Hospital, New York

SECOND EDITION

ILLUSTRATED

ST. LOUIS
THE C. V. MOSBY COMPANY

1949

COPYRIGHT 1937-1949 BY THE C. V. MOSBY COMPANY
(All rights reserved)

First Edition Reprinted
August, 1938

Printed in the
United States of America

Branch of
The C. V. Mosby Company
St. Louis

PREFACE TO SECOND EDITION

Work upon a new and enlarged edition of Kantor's *Synopsis of Digestive Diseases* was just getting under way at the time of his death and as his close associate the task of completing the work fell to me. On approaching the problem however it soon became apparent that because of the recent accumulation of new and important information resulting from research in gastroenterology a book of the synopsis type would be inadequate. It was decided therefore entirely to rewrite the book to reset it in a new format and to publish it as a *Handbook of Digestive Diseases*.

This volume represents an attempt to put down the salient and essential facts concerning gastrointestinal diseases. It is by no means intended to be a comprehensive treatise. Rather it is a twofold effort first a concise presentation of the fundamental aspects of digestive diseases and second an approach to the subject on a sound physiological basis. The fact that gastroenterology is an important and inseparable segment of internal medicine has been stressed throughout.

The close relationship I was privileged to enjoy with Dr. Kantor in private practice in clinical work on the wards of Montefiore Hospital and in the teaching of postgraduate students will undoubtedly be reflected in this book. The *Synopsis* was unique in that it essentially represented Dr. Kantor's own viewpoint. This book is based in part on the course in gastroenterology given annually at Montefiore Hospital. In addition I have drawn heavily upon the observations of my colleagues at Bellevue and Montefiore Hospitals. To the latter institution I owe a debt of gratitude for the liberal use granted me of its unmatched collection of clinical and pathological data. A critical

evaluation of many important contributions by others in this field has been included and for those who wish to explore the subject further a selected bibliography largely limited to recent contributions has been added. This book is essentially a new work for which I bear the responsibility.

I should like to thank Dr. Louis Leiter, Chief of the Medical Division, Montefiore Hospital, who read the manuscript and offered many critical and constructive suggestions. My indebtedness to him is profound. In the preparation of the chapters on gastritis and peptic ulcer I have had the invaluable assistance of Dr. Joseph B. Kirsner of the University of Chicago. To Dr. Alfred Angrist I am obligated for assistance in describing the pathological specimens and photomicrographs and to Dr. Harry D. Lem for aid in preparing the chapter on Diseases of the Mouth. Dr. Morris Pearlmuter has critically read and edited a large part of the book. For the opportunities afforded me to study gastrointestinal cases on the wards of the Fourth Medical (New York University) Division, Bellevue Hospital, I am indebted to the Director, Dr. Charles H. Nammack. Miss Margaret Bothyl, Medical Records Librarian, Montefiore Hospital, was extremely helpful in providing data incorporated in the text. My secretary, Miss Lillian Majestic, has worked indefatigably to prepare the manuscript for publication.

The sections on Irritable Colon and Constipation have been taken in large part from Dr. Kantor's contributions on these subjects in *Portis's Diseases of the Digestive System*. I am grateful to Dr. Sidney A. Portis and to the publishers Lea & Febiger for permission to use this material. The same publishers have allowed me to quote from *Drugs Used in Clinical Diagnosis* (Part II), *Review of Recent Literature*, which appeared in the *American Journal of Medical Sciences*, November 1940.

The kind understanding and consideration which my wife gave to the problem at hand greatly eased and helped along an undertaking which otherwise would have been forbidding.

Finally, without the helpful interest and insight through the years of Lavette B. Dow this and much else would have been impossible.

ANTHONY M. KASICH

New York

PREFACE TO FIRST EDITION

This book is an attempt to present simply, clearly and concisely the essential facts concerning the diseases of digestion. An effort has been made to preserve the proper balance between stress and subordination, inclusion and omission and to emphasize throughout the ways in which gastroenterology fits into the larger field of internal medicine. To this end the opening chapter on classification has been designed for early orientation. Clinical syndromes common to many diseases have been elaborated as fully as possible and a final section has been devoted to digestion symptoms in extradigestive diseases.

In the preparation of this book the writer has drawn on his clinical records for much of his basic statistics as well as on his practical experience in teaching gastroenterology to both undergraduate and graduate students. The current literature in periodicals and in special monographs has been carefully and it is hoped critically utilized. Owing to the limitations of space individual acknowledgments cannot be given as freely as the writer would wish or in the form of complete references. Only those names are cited which are required for credit because of literal or approximate quotation.

The illustrations, most of them original and all of them drawn especially for the author by the skilled hand of Mr Alfred Feinberg, include several synoptic charts devised for the diagrammatic presentation of the more important diseases. The sketch of mucosal visualization in Fig. 18 is based on an illustration in Hurst and Stewart's *Gastric and Duodenal Ulcer* and that of volvulus in Fig. 20 is copied from Thorek's article (*J A M A* 81: 639, 1923).

The special diet forms are derived in practically all instances from the Montefiore Hospital Manual of Diets prepared under the guidance of Miss Lenna F. Cooper.

The manuscript has been read by Dr Allen O Whipple Professor of Surgery Columbia University Dr L Lachwitz Chief of Medical Division Montefiore Hospital and Dr Jerome A Marks Visiting Physician Harlem Hospital The Section on schistosomiasis has been read by Dr Ramon J Sifre Assistant Professor of Hygiene in the School of Tropical Medicine Puerto Rico The writer wishes to express his thanks to all these for their courtesy

JOHN L KANTOR

New York

CONTENTS

CHAPTER I

CLASSIFICATION OF DIGESTIVE DISORDERS	PAGE 17
---------------------------------------	---------

CHAPTER II

DIAGNOSTIC METHODS HISTORY AND PHYSICAL EXAMINATION	18
The History 18 The Physical Examination 27	

CHAPTER III

DIAGNOSTIC METHODS SPECIAL TESTS	40
Gastric Analysis 40 Gastric Test Meals 42 Duodenal Drainage 52 Intestinal Test Meals 55 The Feces 56 Proctoscopy 57 The Poentgen Examination 60 Cholecystography 66	

CHAPTER IV

ORGANIC CONSTITUTIONAL INFERIORITY THE ANOMALIES	71
Body Habitus 72 Visceroptosis 74 Diverticulosis 77	

CHAPTER V

DISEASES OF THE MOUTH	79
Disorders of Mastication 79 Disorders of Excretion 80 Infections 81 Oral Manifestations of Deficiency Disorders 82 Neurotic Disorders 90 Disorders of the Salivary System 90	

CHAPTER VI

DISEASES OF THE ESOPHAGUS	93
Anomalies 93 Diverticulum 93 Cardiospasm 95 Peptic Ulcer of the Esophagus 99 Esophagitis 100 Stricture of the Esophagus 101 Hypertical Dysphagia (Plummer Vinson Syndrome) 101 Esophageal Varices 102 Scleroderma 104 Carcinoma of the Esophagus 107 Uncommon Diseases of the Esophagus 111	

CHAPTER VII

DISEASES OF THE STOMACH	114
Anomalies 114 Diaphragmatic Hernia 115 Esophageal Hiatus Diaphragmatic Hernia 116 Congenital Pyloric Stenosis 124 Functional Dyspepsia 125 The Syndrome of Gastric Irritation 128 Standard Bland Diet 129 Achlorhydria and Achylia Gastrica 130 Achlorhydria Associated With Organic Disease Either of the Digestive Tract or of Other Organs 132 Gastritis 134 Chronic Gastritis 136 Acute Dilatation of the Stomach 141	

CHAPTER VIII		PAGE
DISEASES OF THE STOMACH (CONTINUED)		145
Peptic Ulcer	145	Ulcer Diet—Modified Sippy Plan
Convalescent Ulcer Diet	174	General Directions to Be Followed for Life by Patients With Peptic Ulcer
Other Forms of Therapy	183	
CHAPTER IX		
DISEASES OF THE STOMACH (CONTINUED)		191
Complications of Peptic Ulcer	191	Peptic Ulcer in Locations Other Than Stomach or Duodenum
	210	Jejunul Ulcer
	211	
CHAPTER X		
DISEASES OF THE STOMACH (CONTINUED)		217
Cancer of the Stomach	217	Sarcoma of the Stomach
Syphilis of the Stomach	232	Tuberculosis of the Stomach
	233	Volvulus of the Stomach
	234	Foreign Bodies in the Stomach
	236	
CHAPTER XI		
DISEASES OF THE SMALL INTESTINE		241
Anomalies of the Duodenum	241	Diverticulum of the Duodenum
	244	Duodenitis
	246	Carcinoma of the Duodenum
	246	Diverticula of Jejunum and Ileum
Atresia of the Small Intestine	249	Meckel's Diverticulum
	249	Hernia
	251	Fistulas of the Small Intestine
	252	Intestinal Obstruction
	256	Ileus
	256	Intestinal Obstruction
	261	Volvulus
	261	Tumors of the Small Intestine
	262	
CHAPTER XII		
DISEASES OF THE SMALL INTESTINE (CONTINUED)		268
Tuberculosis	268	Mesenteric Lymph Node Tuberculosis (Tabes Mesenterica)
	273	Syphilis
	273	Amyloid Disease
	274	Diarrhea
	275	Enteritis
	280	Regional Ileitis
Idiopathic Steatorrhea (Sprue Celiac Disease)	290	In
testinal Lipodystrophy (Lipophagia Granulomatosis		Whipple's Disease)
	300	
CHAPTER XIII		
DISEASES OF THE COLON		304
Anomalies	304	Congenital Megacolon (Hirschsprung's Disease)
	311	Constipation
	313	The Unstable Colon (Irritable Colon Spastic Colon Simple Colitis)
	322	Mucous Colitis
	328	

CHAPTER XIV

PAGE

DISEASES OF THE COLON (CONTINUED)

332

Bacillary Dysentery 332 Ulcerative Colitis 343 Ulcerative Colitis Diet 358 Intestinal Protein Diet 358 Diverticulosis and Diverticulitis 362 Tumors of the Colon and Rectum 367 Benign Tumors 367 Diffuse Adenomatosis (Hereditary Polyposis) 368 Carcinoma of Colon and Rectum 369 Endometriosis of the Intestine 381

CHAPTER XV

DISEASES OF THE APPENDIX

387

Acute Appendicitis 387 Chronic Appendicitis 390 Tumors of the Appendix 391

CHAPTER XVI

DISEASES OF THE RECTUM AND ANUS

393

Proctitis and Periproctitis 393 Fistula Fissure and Stricture of the Rectum 394 Lymphopathia Venereum (Lymphogranuloma Inguinale) 395 Hemorrhoids 399 Carcinoma 400

CHAPTER XVII

DISEASES OF THE LIVER

401

Anomalies 401 Functions of the Liver and Liver Function Tests 401 Jaundice 414 Obstructive Jaundice 416 Congenital Hemolytic Jaundice 420 Constitutional Hepatic Dysfunction 422

CHAPTER XVIII

DISEASES OF THE LIVER (CONTINUED)

426

Infectious (Epidemic) Hepatitis 426 Chronic Infectious Hepatitis in Women After the Menopause 443 Leptospirosis Icterohaemorrhagica (Weil's Disease Infectious Jaundice Spirochetal Jaundice) 444 Arsenical Hepatitis (Postarsphenamine Jaundice) 448 Affections of the Hepatic Blood Vessels Chronic Passive Congestion of the Liver (Nutmeg Liver Cardiac Cirrhosis) 450 Diseases of the Hepatic Veins Chlari's Syndrome 454

CHAPTER XIX		PAGE
DISEASES OF THE LIVER (CONTINUED)		460
Fatty Metamorphosis of the Liver 460 Portal Cirrhosis (Laennec's Cirrhosis Atrophic Cirrhosis) 462 Hemochromatosis 471 Intrahepatic Biliary Obstruction (Hypertrophic Biliary Cirrhosis) 473 Hepatolenticular Degeneration (Wilson's Disease) 474 Liver Abscess (Suppurative Hepatitis) 475 Tumors of the Liver 477 Benign Tumors 477 Carcinoma of the Liver 477 Sarcoma of the Liver 481 Echinococcus Cysts 482		
CHAPTER XX		
DISEASES OF THE GALLBLADDER AND BILE DUCTS		485
Cholecystitis 485 Cholelithiasis 488 Low Fat Low Cholesterol Diet 495 Stricture of the Bile Ducts 497 Carcinoma of the Bile Ducts 499 Choledocholithiasis 501 Carcinoma of the Gallbladder 503		
CHAPTER XXI		
DISEASES OF THE PANCREAS		506
Anomalies 506 Insufficiency 506 Hyperinsulinism 507 Acute Pancreatitis 508 Chronic Pancreatitis 513 Benign Cysts 516 Pancreatic Lithiasis 520 Tumors of the Islands of Langerhans 522 Carcinoma of the Pancreas 525		
CHAPTER XXII		
INTESTINAL PARASITES THE PROTOZOA		541
Amebiasis 541 Flagellates (Infusoria) Giardia 552		
CHAPTER XXIII		
INTESTINAL PARASITES (CONTINUED) THE FLATWORMS CESTODES		556
Taenia Saginata 556 Taenia Solium 560 Dibothriocephalus Latus 560 Taenia Echinococcus 561		
CHAPTER XXIV		
INTESTINAL PARASITES (CONTINUED) THE FLATWORMS TREMATODES		564
The Blood Flukes (Schistosomes) 564 Schistosomiasis (Bilharziasis) Produced by Schistosoma Mansoni 566 Schistosomiasis Japonica (Oriental Schistosomiasis Katayama Disease) 571 Schistosomiasis Due to Schistosoma Haematobium 577 The Liver Flukes 578 Clonorchis Sinensis 578 Fasciola Hepatica 579 Intestinal Flukes 580		

CHAPTER XXV

PAGE

INTESTINAL PARASITES (CONTINUED) THE ROUNDWORMS 582

Trichina 582 Uncinaria 586 Ascaris 589 Pinworm
590 Whipworm 592 Strongyloides Stercoralis 592

CHAPTER XXVI

DIGESTIVE SYMPTOMS IN EXTRADIGESTIVE DISEASES 595

Nervous Cardiovascular and Respiratory Diseases 595
Nervous Diseases 597 Cardiovascular Diseases 596
Respiratory Diseases 598 Diseases of the Blood Metabolism and the Genitourinary System Plumbism 599
Diseases of the Blood 599 Diseases of the Metabolism
599 Diseases of the Genitourinary System 600 Plumbism
601

CHAPTER XXVII

PSYCHIATRIC ASPECTS OF DIGESTIVE DISEASES

JAMES A. BRUSSEL, M.D. 603

CHAPTER XXVIII

GASTROINTESTINAL ALLERGY HARRY SWARTZ, M.D. 618

HANDBOOK OF DIGESTIVE DISEASES

Chapter I

CLASSIFICATION OF DIGESTIVE DISORDERS

For purposes of discussion digestive disorders may be grouped under one or more of the following heads

- 1 Those due to constitutional digestive inferiority
- 2 Those due to acquired digestive disease
- 3 Those due to extradiigestive disease

Constitutional inferiority may in turn be divided into organic and functional types. Organic constitutional inferiority results from congenital anomalies of the digestive tract. Functional constitutional inferiority results from congenital instability of the autonomic nervous system.

Acquired disease is either toxic, metabolic, infectious, traumatic or parasitic in nature. In many cases its origin is obscure.

Primary disease in other systems of the body often causes reflex symptoms in the digestive apparatus. Prominent examples are the dyspepsias of cardiovascular degeneration, of phthisis, of lead poisoning, and of syphilis.

The cause of neoplasms is still unknown, but there seems to be increasing evidence (Mielin) that the predisposition to tumor formation is inherited. If this theory proves to be correct, our conception of the role played by constitutional (i.e. hereditary) factors in the production of digestive disorders will have to be greatly expanded.

CHAPTER III—continued

PAGE

3 33	High voltage sharply peaked T waves in uræmia associated with a high blood potassium The long Q T interval is due to hypocalcæmia	104
3 34	Effect of potassium on the T waves in a case of concordant left ventricular preponderance	105

CHAPTER IV

4 01	Anatomy of the conducting system	108
4 02	Sinus arrhythmia	109
4 03	Sinus tachycardia showed by carotid sinus compression	111
4 04	Relative cardiac enlargement due to sinus bradycardia	113
4 05	Nodal escape in sinus bradycardia	114
4 06	Sino auricular block irregular dropped beats	114
4 07	Sino auricular block rate doubles on effort	115
4 08	Cardiac standstill occurring spontaneously in sino auricular block	115
4 09	Cardiac standstill due to carotid sinus compression	115
4 10	Nodal rhythm—three types (a) (b) and (c)	116
4 11	Shifting nodal rhythm	117
4 12	Prolonged R R interval with P coinciding with the previous T wave	118
4 13	Partial heart block with dropped beats (Wenckebach type)	119
4 14	2-1 heart block	119
4 15	Complete heart block Two examples (a) and (b)	1 1
4 16	Complete A V dissociation with the ventricles beating faster than the auricles	121
4 17	Stokes Adams attack produced by carotid sinus compression in a patient with paroxysmal complete heart block	1-3
4 18	Alternating left bundle branch block	126
4 19	Auricular ectopic beats	127
4 20	Nodal ectopic beats Slight deformity of QRS is due to fatigue block	128
4 21	Right ventricular ectopic beats	1 8
4 2	Left ventricular ectopic beats causing coupling	129
4 23	Interpolated ventricular ectopic beats (lead 3)	131
4 24	Paroxysmal auricular tachycardia terminated by means of meclozin	13-
4 25	Paroxysmal auricular tachycardia blocked by carotid sinus compression	133
4 26	(a) and (b) Two cases of paroxysmal auricular tachycardia showing varying degrees of spontaneous A V block	134
4 27	(a) and (b) Two cases of paroxysmal auricular tachycardia slowed by by means of quinidine	135
4 28	Paroxysmal auricular tachycardia followed by auricular ectopic beats	135
4 9	(a) Paroxysmal nodal tachycardia	136
	(b) Coronary sinus rhythm	136
4 30	Paroxysmal ventricular tachycardia	137
4 31	Auricular flutter with left bundle branch block	137
4 32	Pre excitation (a) Usual appearances (b) Alternating	139
4 33	Wolff Parkinson White syndrome showing paroxysmal tachycardia of two types (a) and (b) followed by auricular re entry (c)	140
4 34	Auricular flutter (a) with 4-1 A V block (b) with 2-1 A V block (c) clarified by means of carotid sinus compression	143
4 35	Auricular flutter treated with digitalis	144
4 36	Lead V ₁ showing coarse auricular fibrillation or impure flutter	146
4 37	Auricular fibrillation	146
4 38	Auricular fibrillation with complete A V dissociation due to digitalis	146
4 39	Auricular fibrillation treated with quinidine	149
4 40	Ventricular fibrillation causing sudden death in a case of ischæmic heart disease	150

CHAPTER V

PAGE

5 01	Relationship of cardiac output to venous filling pressure - Starling's curve	155
5 02	Graph illustrating various physiological changes in an attack of paroxysmal cardiac dyspnoea	159
5 03	Pulmonary congestion in left ventricular failure	161
5 04	Acute pulmonary oedema (a) in left ventricular failure (b) in mitral stenosis	162
5 05	Phonocardiogram showing a normal third heart sound	164
5 06	Electrical alternation in a case of malignant disease involving the pericardium	166
5 07	Photograph showing distension of the external jugular vein in a case of congestive heart failure	169
5 08	Tracings of serial skiagrams of liver and spleen opacified by means of thorotrast to illustrate the effect of digoxin in heart failure	171
5 09	Dependent oedema in congestive heart failure	172
5 10	Fall in erythrocyte sedimentation rate resulting from the development of congestive failure in a case of active rheumatic carditis	175
5 11	Increase in transverse diameter of heart caused by congestive failure	176
5 12	Typical effect of digitalis on venous pressure or right auricular pressure in four cases of congestive heart failure	180
5 13	Typical effect of digitalis on the blood pressure pulse rate and cardiac output in a case of hypertensive heart failure	180
5 14	Graph showing the pronounced slowing effect of digitalis in a case of congestive failure with normal rhythm due to active rheumatic carditis	181
5 15	The action of digitalis on the arm to tongue circulation time and on pulse rate in four cases of left ventricular failure with normal rhythm	182
5 16	Chart showing considerable diuresis resulting from the administration of digitalis to a case of hypertensive heart failure with normal rhythm	182
5 17	Chart illustrating the failure of digitalis to lower the right auricular pressure in twelve cases in which it was raised from causes other than congestive failure	183
5 18	Chart illustrating the beneficial effect of mercurial diuretics in preventing paroxysmal cardiac dyspnoea	185

CHAPTER VI

6 01	Carotid sinus pressure causing cardiac standstill	196
------	---	-----

CHAPTER VII

7 01	Mirror image dextrocardia. Electrocardiogram showing reversal of all complexes in lead I while leads II and III are interchanged	204
7 02	Unexplained cardiac enlargement in a relatively young man	205
7 03	(a) The six primitive aortic arches (b) Subsequent arrangement of the six primitive aortic arches in man	206
7 04	Diagrams illustrating the three main types of coarctation of the aorta	207
7 05	Notching of the inferior border of the ribs due to pressure erosion from enlarged intercostal arteries in coarctation of the aorta	209
7 06	Visualisation of coarctation of the aorta by means of angiocardio-graphy	209
7 07	(a) Coarctation of the aorta associated with patent interventricular septum proved at necropsy (b) Similar case but without necropsy proof	210
7 08	(a) Catheter in the left ventricular via a patent foramen ovale (b) Catheter in right pulmonary vein via a patent foramen ovale	214
7 09	Catheter in left atrial appendage	214

CHAPTER VII — *continued*

PAGE

7 10	Case of arachnodactyly (a) Facies (b) High arched palate and deformed teeth (c) Spider fingers	215
7 11	Skiagram of a case of atrial septal defect showing gross dilatation of the pulmonary artery and its branches enlargement of the right auricle and hypoplasia of the aorta	217
7 12	Lutembacher's syndrome (a) A P views (b) First oblique position showing dilatation of the left auricle	217
7 13	Atrial septal defect in a child aged ten	218
7 14	Electrocardiogram in a case of atrial septal defect showing right bundle branch block	219
7 15	Functional studies in atrial septal defect	219
7 16	Maladie de Roger—X ray appearances	221
7 17	Skiagram of a case of ventricular septal defect with considerable increase of pulmonary blood flow	221
7 18	Skiagram of a case of patent ductus showing enlargement of left ventricle but little dilation of the pulmonary artery	224
7 19	Skiagram of a more advanced case of patent ductus showing considerable left ventricular enlargement and engorgement of the pulmonary vessels in addition to dilatation of the pulmonary artery	225
7 20	Electrocardiogram in a case of patent ductus showing a strong QR pattern and inverted U waves in lead V ₅	226
7 21	Electrocardiogram in a case of patent ductus showing gross left ventricular preponderance	226
7 22	Average catheter findings in patent ductus	227
7 23	Case of patent ductus arteriosus (a) before and (b) one year after successful ligation	228
7 24	Skiagram of a case of simple pulmonary stenosis showing dilatation of the pulmonary artery and hypoplasia of the aorta	230
7 25	Electrocardiogram of a case of simple pulmonary stenosis showing right ventricular preponderance	230
7 26	Pulmonary valvular stenosis with reversed interatrial shunt showing (a) dilatation of the pulmonary arc alone (b) considerable cardiac enlargement	233
7 27	Strong ventricular dominance due to pulmonary valvular stenosis with reversed interatrial shunt	233
7 28	Findings on cardiac catheterisation in a typical case of pulmonary valvular stenosis with reversed interatrial shunt	234
7 29	Catheter penetrating the left auricle through an atrial septal defect showing the high position of the latter	234
7 30	Catheter lying in the pulmonary veins (a) Right upper (b) Right lower (c) Left upper	235
7 31	Skiagram of a case of Fallot's tetralogy showing the cœur en sabot (a) A P view (b) Second oblique position	237
7 32	Right sided aortic arch in a case of Fallot's tetralogy	237
7 33	Angiocardiogram of a case of Fallot's tetralogy showing simultaneous opacification of the aorta and pulmonary artery	238
7 34	Electrocardiogram of a case of Fallot's tetralogy showing marked right ventricular dominance	238
7 35	Catheterisation of the pulmonary artery in a case of Fallot's tetralogy (a) Main trunk (b) Right branch (c) Left branch	239
7 36	Average catheter findings in Fallot's tetralogy	240
7 37	Penetration of the foramen ovale and pulmonary veins in a case of Fallot's tetralogy	240
7 38	Site of abrupt pressure change in pulmonary valvular stenosis	240
7 39	Pulmonary subvalvular stenosis the tip of the catheter is still in a low pressure zone	241
7 40	Demonstration of the anatomy of the pulmonary arteries in a case of Fallot's tetralogy (a) Catheter in right pulmonary artery (b) Diodone in left pulmonary artery	241
7 41	Eisenmenger's complex	245

CHAPTER VII —continued

PAGE

7 4	Angiocardiogram in Eisenmenger's complex	245
7 43	Skiagram of a case of transposition of the great vessels associated with atrial and ventricular septal defects	47
7 44	Skiagram of a case of transposition of the great vessels showing penetration of the pulmonary artery from the right ventricle via the ventricular septal defect	48
7 45	Skiagram of a case of transposition of the great vessels showing a catheter penetrating the aorta directly from the right ventricle	249
7 46	(a)-(d) Angiocardiographic series in tricuspid atresia	251

CHAPTER VIII

8 01	Macroscopic nodules on the surface of the heart	258
8 02	(a) and (b) Erythema marginatum	262
8 03	Erythema multiforme	263
8 04	(a) and (b) Subcutaneous rheumatic nodules	264
8 05	Skiagram showing rheumatic pneumonia in a girl	265
8 06	(a) (b) and (c) Serial skiagrams showing rapid development of cardiac enlargement as a result of active rheumatic carditis	67
8 07	Electrocardiogram showing prolongation of the PR interval in a case of active rheumatic carditis	270
8 08	Graph showing QT plotted against the sedimentation rate in 60 cases of active rheumatic carditis and in 14 rheumatic fever controls without carditis	271
8 09	Behaviour of QT in six cases of acute rheumatic carditis with rapid clinical recovery	27
8 10	Prolonged QT _c following a recurrence of active rheumatic carditis	27
8 11	Graph showing relapse due to letting up when QT _c was still grossly prolonged	27

CHAPTER IX

9 01	Phonocardiogram showing an innocent systolic murmur at the mitral area	81
9 02	Phonocardiograms in two cases of organic mitral incompetence (a) Showing a systolic murmur (b) Showing a late systolic murmur	83
9 03	Skiagram showing dilatation of the left ventricle in a case of organic mitral incompetence. On fluoroscopy the outline of the left auricle expanded visibly during systole	284
9 04	Phonocardiogram showing a mitral diastolic murmur following the third heart sound in a case of mitral stenosis	286
9 05	Phonocardiogram in a case of patent ductus showing a functional mitral diastolic murmur	287
9 06	Phonocardiogram showing a crescendo presystolic murmur timed against the electrocardiogram and phlebogram	287
9 07	Skiagram of a case of mitral stenosis showing dilatation of the left auricle between the pulmonary arc and the left ventricle	88
9 08	Skiagram of a case of mitral stenosis showing dilatation of the left auricle in the right anterior oblique position. The œsophagus is outlined with barium	289
9 09	Skiagram of a case of mitral stenosis showing dilatation of the left auricle in the left anterior oblique position	289
9 10	Skiagram of a case of mitral stenosis showing miliary nodules in the lungs due to hæmosiderosis	289
9 11	Angiocardiogram in a case of mitral stenosis	289
9 12	Electrocardiogram in a case of mitral stenosis showing widened bifid P waves particularly in leads I, aV ₅ and V ₆ . The heart is vertical	290
9 13	Electrocardiograms showing the P waves in eighteen unselected cases of mitral stenosis	291

CHAPTER IX—continued

PAGE

9 14	Standard lead electrocardiograms in four cases of mitral stenosis showing tall sharp P waves like those seen in pulmonary heart disease	291
9 15	Electrocardiogram in a case of mitral stenosis showing partial right bundle branch block	292
9 16	Phonocardiogram illustrating a diminuendo aortic diastolic murmur	293
9 17	Arteriogram illustrating the water hammer pulse of aortic incompetence	296
9 18	Skigram showing prominence of the aortic arch and enlargement of the left ventricle in a case of aortic incompetence	297
9 19	Skigram in the second oblique position showing unfolding of the aortic arch and enlargement of the left ventricle	297
9 20	Electrocardiogram in a case of aortic incompetence showing evidence of left ventricular enlargement	298
9 21	Arteriogram in a case of aortic stenosis. The percussion wave is prolonged and the maximum pressure is reached late in systole	300
9 22	Arteriogram illustrating pulsus bisferiens in a case of combined aortic stenosis and incompetence	300
9 23	Skigram of a case of aortic stenosis showing great enlargement of the left ventricle slight prominence of the ascending aorta and pulmonary congestion	301
9 24	Electrocardiogram in a case of aortic stenosis showing concordant left ventricular preponderance in standard leads the heart being vertical	30
9 25	Jugular phlebogram showing fusion of the c and t waves in a case of tricuspid incompetence owing to auricular fibrillation the a wave is absent	304
9 26	Skigram showing gross dilatation of the right auricle with a blunt right cardio phrenic angle in a case of tricuspid incompetence	305
9 27	Graph illustrating a fall in mean central venous pressure as the catheter is withdrawn from right auricle and superior vena cava into the sub clavian vein	306
9 28	Electrocardiogram showing exceptionally tall P waves in a case of mitral and tricuspid stenosis	307

CHAPTER X

10 01	(a) and (b) Electrocardiogram in a case of toxic myocarditis due to pneumonia	316
10 02	(a) and (b) Focal necrosis in a case of Fiedler's carditis	319
10 03	Skigram showing general enlargement of the heart in a case of Fiedler's carditis	320
10 04	Electrocardiogram showing coupling due to digitalis	3-3
10 05	Electrocardiogram showing partial heart block due to digitalis	323
10 06	Electrocardiogram showing paroxysmal tachycardia due to digitalis	3 4
10 07	Electrocardiogram showing depression of the RS T segment due to digitalis	324
10 08	Shortening of the Q T interval due to digitalis	3-5
10 09	Widening of the QRS complex and accentuation of the T wave due to a high blood potassium in a case of uremia	3 6
10 10	Graph illustrating a high right auricular pressure that does not respond to digitalis in a case of acute nephritis	3-7

CHAPTER XII

12 01	Electrocardiogram showing the early phase of the pericardial T ₁ pattern	34
12 02	Electrocardiogram showing the later phase of the pericardial T ₂ pattern case of pyogenic pericarditis secondary to broncho pneumonia	343

CHAPTER XII—*continue*

PAGE

12 03	Electrocardiogram showing late changes due to pericarditis	344
12 04	Skiagram of a case of pericardial effusion showing an acute right cardio phrenic angle	345
12 05	Skiagram showing rapid diminution in size of cardiac silhouette as effusion is absorbed	346
12 06	Skiagram showing the blunt right cardio phrenic angle in tricuspid incompetence	347
12 07	Skiagrams of a case of constrictive pericarditis (a) Triangular shaped heart (b) Calcified pericardium	350
12 08	Electrocardiogram in a case of chronic constrictive pericarditis showing low voltage and flat or inverted T waves	350
12 09	Pericardial effusion of three years duration in a case of extreme essential hypertension	355

CHAPTER XIII

13 01	Saccular aneurysm of the ascending aorta	359
13 02	Angiocardiogram showing partial superior vena cava obstruction due to an aneurysm	360
13 03	Skiagram showing several aneurysms of the aortic arch (a) A P view (b) Second oblique position	361
13 04	Skiagram showing erosion of the bodies of several dorsal vertebræ as the result of pressure from an aneurysm	362
13 05	Angiocardiogram outlining a normal pulmonary artery (a) and aorta (b) in a case of mediastinal tumour (c)	362
13 06	Skiagram of a case of syphilitic aortic incompetence showing fusiform dilatation of the ascending aorta and gross enlargement of the left ventricle (a) A P view (b) Left anterior oblique position	365

CHAPTER XIV

14 01	(a) Skiagram of the heart in a case of occlusive coronary atherosclerosis the coronary vessels have been injected with a radio opaque gel (b) Normal control for comparison	372 373
14 02	Diagram illustrating radiation of pain in ischæmic heart disease	375
14 03	Electrocardiogram (a) before and (b) after exertion sufficient to cause breathlessness and fatigue in a case subject to attacks of angina pectoris the control record (a) is normal the second record (b) shows significant depression of the S T segment in practically all leads	379
14 04	Graph showing close correlation between the height of the blood pressure and the degree or extent of pain during an attack of angina pectoris treated with trinitrin	381
14 05	Graph illustrating rapid correction of ischæmic depression of the S T segment in lead V ₃ in a patient with angina pectoris by means of trinitrin	382
14 06	Electrocardiogram during an attack of myocardial ischæmia treated with amyl nitrite—expected response	383
14 07	Paradoxical effect of amyl nitrite on the S T segment	384
14 08	Chart illustrating the prevention of cardiac infarction by means of anticoagulants angina at rest continued for two months	385
14 09	Behaviour of the blood pressure in four cases of acute myocardial infarction	390
14 10	Behaviour of the sedimentation rate in four cases of acute myocardial infarction	390
14 11	Electrocardiogram showing antero lateral cardiac infarction Maximum changes are seen in leads V ₅ and V ₆ VL and lead I	391
14 12	Electrocardiogram showing antero septal cardiac infarction Maximum changes are seen in leads V ₃ and V ₄	392

CHAPTER XIV—continued

	PAGE
14 13 (a) Electrocardiogram in posterior cardiac infarction (a) Characteristic changes are seen in lead VF and hence in leads 2 and 3 The ST segment is depressed in lead V ₄ (b) Later stage	393
14 14 (a) (b) and (c) Electrocardiogram showing widespread monophasic Q waves and persistent elevation of the S T segment associated with ventricular aneurysm	394
14 15 Electrocardiogram of a case of old cardiac infarction showing persistent Q waves and Pardee coving of the ST segment in anterior left ventricular surface leads and their counterparts (leads VL and standard lead I) The infarct occurred fourteen months previously	395
14 16 Electrocardiogram in a case of pregnancy showing a prominent Q wave and inversion of the T wave in lead 3 due to cardiac rotation note the absence of a pathological Q wave in lead VF and the presence of an S wave in standard lead I	395
14 17 Electrocardiogram showing typical appearances of anterior cardiac infarction in the presence of right bundle branch block	396
14 18 Electrocardiogram showing typical appearances of anterior cardiac infarction in the presence of left bundle branch block	396
14 19 Skiagram in a case of anterior cardiac infarction showing a ledge on the left border of the heart	398
14 20 Kymogram of a case of anterior cardiac infarction showing an area of absent pulsation on the left border of the heart near the apex	399
14 21 Kymogram of a case of hypertensive heart failure showing absence of pulsation at the apex	400
14 22 Skiagram of a case of ventricular aneurysm	401
14 23 Skiagram of a case of organic mitral incompetence showing a dilated left auricle on the left border of the heart	40*
14 24 Skiagram of a case of stab wound of the heart showing a hæmatoma on its left border (successfully evacuated later)	403
14 25 Electrocardiogram showing the mode of death in a case of ischæmic heart disease ventricular fibrillation developed while a routine graph was being taken	403
14 26 Cardiac infarction complicated by perforation and hæmopericardium (a) Original anterior infarct (b) After recovery (c) After perforation	405
14 27 Electrocardiogram showing persistent depression of the ST segment in a case of acute coronary insufficiency	409
14 28 Electrocardiogram showing transient inversion of the T waves following prolonged circulatory collapse with extreme tachycardia without evidence of structural disease of the heart	410
14 29 (a) and (b) Transient inversion of the T waves due to carbon monoxide poisoning	411

CHAPTER XV

15 01 Electrocardiogram in a case of hypertensive heart disease (see text)	4 6
15 02 Electrocardiogram in a case of hypertensive heart disease with clockwise rotation about the longitudinal axis the transition zone is shifted to the left	4 7
15 03 Electrocardiogram in a case of hypertensive heart disease with anti-clockwise rotation about the longitudinal axis the transition zone is shifted to the right	427
15 04 Electrocardiogram in a case of hypertensive heart disease The heart is electrically vertical	428
15 05 Electrocardiogram showing concordant left ventricular preponderance due to a semi vertical position of the heart	428
15 06 Hypertensive heart disease showing left ventricular enlargement (a) A P view (b) Angiocardiogram in second oblique position	430
15 07 Skiagram of a case of hypertensive heart disease showing unfolding of the aortic arch (a) A P view (b) Second oblique position (c) First oblique position	431

CHAPTER XV —continued

PAGE

- 15 08 Right anterior oblique view of a case of hypertensive heart disease showing backward displacement of the oesophagus at left auricular level 431
- 15 09 Comparison of the coronary systems in a normal (a) and a hypertensive heart (b) 432
- 15 10 Coronary systems of two cases of hypertensive heart disease with angina pectoris 43

CHAPTER XVI

- 16 01 Behaviour of the blood pressure in nine cases of massive pulmonary embolism 446
- 16 02 Behaviour of the venous pressure in eight cases of massive pulmonary embolism 448
- 16 03 Electrocardiograms showing the characteristic appearances associated with massive pulmonary embolism 450
- 16 04 Electrocardiogram showing transient right bundle branch block in a case of massive pulmonary embolism 451
- 16 05 Skiagram showing a small pulmonary infarct at the right base with a little hæmorrhagic effusion 452
- 16 06 Effect of a venous pressure lowering agent on the blood pressure and cardiac output of a case of massive pulmonary embolism 456
- 16 07 Skiagram showing miliary embolic carcinomatosis of the lungs 458
- 16 08 Radiological appearances of the lungs showing embolic secondaries due to chorion epithelioma 458

CHAPTER XVII

- 17 01 Electrocardiogram showing prominent P waves and right ventricular dominance in a case of idiopathic pulmonary hypertension 463
- 17 02 Skiagram showing prominence of the pulmonary arc and right ventricular enlargement in a case of idiopathic pulmonary hypertension 463
- 17 03 Electrocardiogram in a case of emphysema showing a vertical electrical position and clockwise rotation (viewed from below) 466
- 17 04 Standard lead electrocardiographic findings in 100 cases of anoxic pulmonary heart disease 466
- 17 05 Chest lead findings in anoxic pulmonary heart disease 467
- 17 06 Standard limb lead electrocardiograms of 32 unselected cases of anoxic pulmonary heart disease with relatively low voltage showing the frequency, amplitude and shape of the pulmonary wave 468
- 17 07 Standard limb lead electrocardiograms of a further 16 unselected cases of anoxic pulmonary heart disease with normal voltage showing the frequency, amplitude and shape of the pulmonary P wave 469
- 17 08 The maximum P waves in one or other of the standard leads (usually lead Δ_2) of 72 unselected normal controls are shown for comparison with figs 17 06 and 17 07 470
- 17 09 (a) and (b) Skiagrams of two advanced cases of anoxic pulmonary heart disease showing dilatation of the pulmonary arc and of the left and right branches 471
- 17 10 (a) Right anterior oblique position showing the increased density and diameter of the pulmonary artery at its bifurcation (b) Left anterior oblique position showing the left pulmonary artery forming an arc almost as dense and as large as the aortic arch 471
- 17 11 Development of aneurysmal dilatation of the right pulmonary artery in a case of anoxic cor pulmonale 475

CHAPTER XVIII

PAGE

18 01	(a) and (b) Exophthalmic goitre	482
18 02	Unilateral lid retraction and exophthalmos	483
18 03	Lid retraction and characteristic thyrotoxic stare	484
18 04	Substernal goitre revealed by X rays	485
18 05	Skiagram showing slight prominence of the aortic knuckle and of the left pulmonary arc in a case of thyrotoxicosis	487
18 06	Electrocardiograms showing relatively high voltage P and QRS waves in six cases of thyrotoxicosis	488
18 07	Effect of thiouracil on the excretion of creatine in the urine	490
18 08	Skiagram showing gross cardiac enlargement in a case of thyrotoxicosis plus mitral stenosis	494
18 09	(a) Thyrotoxic heart failure (b) After subtotal thyroidectomy	495
18 10	(a) Electrocardiogram showing sinus bradycardia low voltage auricular and ventricular complexes and flat T waves in all leads in a case of myxœdema (b) Normal electrocardiogram after treatment	497
18 11	Electrocardiogram before and after treatment in a case of cretinism	497
18 12	(a) Myxœdema (b) After seven weeks treatment	498
18 13	(a) Skiagram showing general enlargement of heart in a case of myxœdema (b) After treatment	498

CHAPTER XIX

19 01	(a) Electrocardiogram showing low voltage and flat or inverted T wave in all leads in a case of pernicious anemia (b) After treatment	504
19 02	Electrocardiogram showing depression of the ST segment due to acute coronary insufficiency resulting from post hæmorrhagic anemia	505
19 03	Skiagram showing general cardiac enlargement in a case of severe pernicious anemia (a) Before and (b) after treatment of the anemia	505
19 04	Electrocardiogram showing the characteristic appearances associated with pregnancy	507
19 05	(a) Skiagram showing a congenital arterio venous aneurysm of the lung (b) Angiocardiogram showing diiodone filling the aneurysm	511
19 06	Calcification in the wall of an arterio venous aneurysm	512

CHAPTER XX

20 01	Case of ruptured mycotic aneurysm of the sinus of valsalva into the pulmonary artery (a) A P view (b) Second oblique position	524
20 02	Skiagram showing machine gun bullet embedded in the wall of the right auricle	526
20 03	Skiagram showing machine gun bullet embedded in the heart since 1917 (see text)	5 6
20 04	Localised pericardial hæmatoma superficially resembling a cardiac aneurysm	527

CHAPTER XXI

21 01	Classical facies build and posture of a case of Da Costa's syndrome Painted by Ian Tizard (Life sized portrait in the museum of the Post graduate Medical School of London)	539
-------	---	-----

•

DISEASES OF THE HEART
AND CIRCULATION

CHAPTER I

APPROACH TO CARDIOLOGY

H EART disease is by far the most common cause of natural death in civilised communities in the more temperate zones of the world. It is responsible for about one third of all such deaths and for an annual mortality rate in the general population of about 0.4 per cent. Both incidence and mortality curves have been rising steadily for many years, a fact which is not fully explained by ageing populations and by the control of infectious fevers, pulmonary tuberculosis and pyogenic infections. The incidence and mortality rate of cancer, for example, shows no comparable rise. Ischæmic heart disease, particularly, is on the increase.

The relative incidence of the various forms of heart disease classified according to etiology is given below.

	<i>Per cent</i>
Congenital heart disease	2
Rheumatic heart disease	25
Bacterial endocarditis	2
Syphilitic aortitis	3
Ischæmic heart disease	25
Hypertensive heart disease	30
Pulmonary heart disease	5
Thyrotoxic heart disease	5
Miscellaneous and uncertain	3
	<hr/>
	100
	<hr/>

HISTORY TAKING

To take an accurate and relevant history is one of the most difficult and important arts in medicine. Sometimes a complete diagnosis can be made from the history alone, and not infrequently the possibilities can be whittled down to two or three. A good history should at least indicate the system involved, or it should point unerringly to some group or groups of diseases. A common mistake is the failure to analyse any given symptom sufficiently. In cardiovascular work this applies especially to pain, breathlessness, palpitations and syncope. The student is usually taught to encourage the patient to tell his story in his own words, and to record them more or less verbatim. Yet such an account may be verbose, irrelevant, inaccurate and misleading. It is an axiom that the leading question must be avoided at all cost, yet again an experienced physician must know that

the ability to put the appropriate leading question at the right moment and the intelligent interpretation of its reply are invaluable. It is not pretended that leading questions may not lead to false information if the power of their suggestion is not appreciated by the questioner and it is agreed that much may be lost by failure to allow the patient freedom and time to express his complaints in his own way but the average patient will not mention half the available information until he is pressed and the data freely given must be checked as at the bar. For example in the differential diagnosis between a neural and non neural somatic lesion an accurate description of the quality of the pain may determine the issue immediately yet the majority of patients will volunteer no information concerning the quality of pain and if asked to describe it will do so inadequately. They may say it is aching or sharp but fail to enlarge on this even when urged to do so. In answer to the leading question 'Does it tingle?' however they may reply at once in the affirmative. It is essential to realise that the matter does not end there that such a positive reply to a leading question demands the most penetrating cross examination until the questioner is satisfied that the pain really does tingle and that the patient is not merely saying so because it seems the easier answer. It is scarcely too much to say that the best history taker is he who can best interpret the answer to a leading question. Appropriate leading questions can only be asked however when the proffered history has provided sufficient data upon which to work and if the physician has sufficient knowledge of the possibilities then entailed. It is this latter factor which makes it easier for the expert than for the student.

CLINICAL EXAMINATION

There are two methods of examining a patient the first begins at the top of the head and ends with the toes a method often adopted for the sake of convenience the second is to examine the various systems of the body one by one in logical sequence. The procedure recommended here is concerned only with the cardiovascular system but it is essential of course that all other systems be examined.

Inspection While extracting the history the physician should be making a preliminary general inspection. He should pay particular attention to the head and neck looking for goitre and for the eye signs of thyrotoxicosis for Corrigan's sign and especially for jugular pulsation. He will note the general build and appearance of the patient his attitude and demeanour and should form some idea of his character. He should observe plethora pallor or cyanosis. He may see that respiration is hurried irregular shallow or wheezy or he may detect the tell tale sigh of emotional tension. He is sure to glance at the hands noting their posture shape colour and behaviour he may discern clubbing of the fingers spooning of the nails tremor or palmar sweating. All these things and many others he will learn to observe without effort taking note of them without seeming to do so and

in such a limited survey may be put on the track of the correct diagnosis and be forewarned where to look most diligently for further signs

Determining the presence or absence of congestive heart failure Congestive heart failure is revealed by an elevated systemic venous pressure distension and tenderness of the liver and dependent dropsy Left ventricular failure on the other hand is more clearly indicated by the history The venous pressure may be assessed by inspecting the cervical veins and is fully considered in Chapter V The examiner should not feel guilty if he switches from inspection of the neck to palpation of the liver to discover whether or not that organ is engorged nor should he feel embarrassed if he desires to turn from the right hypochondrium to the feet and sacral region in search of œdema if he be criticised for gymnastics he may reply that he is more concerned with the logical sequence of his thoughts

The pulse It is customary to examine the pulse first at the wrist and to consider it in terms of speed rhythm tension amplitude and quality at the same time it is convenient to note the state of the arterial wall Whilst speed and rhythm may be checked by auscultation of the heart and tension by sphygmomanometry the quality and amplitude of the pulse wave can only be analysed in peripheral vessels and are features of great diagnostic importance Thus an anacrotic or plateau pulse of small volume signifies aortic stenosis a bisferiens pulse combined aortic stenosis and incompetence a water hammer pulse vasodilatation and so on Attention should next be directed to the other radial artery thence to the brachials carotids femorals posterior tibials and finally to the dorsal arteries of the feet Difficulty in locating the radial artery may be due to its taking an aberrant dorso lateral course Weakness on one or other side usually denotes proximal compression as from aneurysm of the aorta but a weak left radial pulse may be due to an ectopic origin and aberrant course of the left subclavian artery Examination of the brachial arteries is particularly fruitful both with respect to the pulse wave and to the state of the vessel itself and should never be neglected The carotids may present a thrill or shudder suggesting aortic stenosis Corrigan's sign indicating aortic incompetence or kinking from atherosclerosis Routine palpation of the femorals would insure the immediate recognition of coarctation of the aorta in nearly all cases diminished and delayed pulsation being characteristic and pathognomonic The presence of pulsation in the vessels of the feet should always be recorded if only for subsequent reference Finally the colour and temperature of the hands and feet should be noted if they are warm an attempt should be made to detect digital throbbing and capillary pulsation The latter is best demonstrated by means of transillumination

The blood pressure Approximate estimation of the blood pressure by clinical means is not only possible but should be practised regularly with experience it is easy to tell whether it is low normal or high and the procedure takes but a moment The physician should stand in front and to the right of the patient and should compress the right brachial artery with his

right thumb while feeling the right radial pulse with the fingers of his left hand the force required to obliterate the pulse represents the systolic blood pressure The alternative method of placing three fingers on the radial artery the first to compress the vessel above the second to feel the pulse and the third to obliterate the ulnar collateral below is difficult cumbersome and less reliable

In cardiovascular work however the blood pressure should always be measured with a mercurial manometer or if an aneroid instrument is used it should be calibrated at frequent intervals against the standard mercurial type The patient must be comfortable whether lying or sitting and must have had time to recover from any recent excitement or exertion The arm should be bared to the shoulder to avoid constriction from clothing and to facilitate proper application of the cuff The latter should be fitted closely and evenly round the arm so that its lower edge is one inch above the bend of the elbow and the middle of the rubber bag lies over the brachial artery Preliminary readings should be taken by palpation as the cuff is inflated the point at which pulsation can no longer be felt in the brachial artery represents the systolic blood pressure as the cuff is deflated brachial pulsation gradually assumes a water hammer quality and then abruptly resumes its normal character the reading corresponding to this sudden change represents the diastolic blood pressure Readings obtained on inflation should be checked on deflation and vice versa When approaching an end point the pressure must be altered slowly in the cuff The palpatory method avoids the pitfall of the auscultatory gap, and is uninfluenced by subjective auditory defects nevertheless it must be checked by auscultation The stethoscope should be applied lightly and accurately over the brachial artery just below but not in contact with the cuff The latter is then inflated to a pressure of some 30 mm Hg above the systolic pressure as found by palpation and slowly deflated The accepted systolic blood pressure is the highest level at which successive sounds are heard As the pressure is further lowered in the cuff the dull thud of the upper limits is replaced first by a murmur and then by louder and sharper sounds the point at which these slapping sounds suddenly become muffled should be taken as the diastolic pressure When there is vasodilatation especially when associated with aortic incompetence sounds may still be heard when the cuff pressure is reduced to zero but normally they disappear a few mm Hg below the change over

When there is hypertension sounds may disappear as the cuff is inflated but reappear at higher levels This is the auscultatory silent gap and though uncommon is not rare preliminary palpation prevents possible error If there are ectopic beats the higher pressure of the beat which follows the ectopic should be ignored In auricular fibrillation only approximate readings can be obtained the systolic pressure should be taken at the point where the majority of beats come through the diastolic where the majority of beats become muffled As the blood pressure normally varies by a few

mm Hg with respiration it may be suitably recorded to the nearest multiple of five

The above recommendations are freely borrowed from the joint report of the committees appointed by the British Cardiac Society and the American Heart Association for the standardisation of methods of measuring the arterial blood pressure (1939)

The normal systolic blood pressure lies between 95 and 145 mm Hg. Whilst it is true that apparently normal subjects between the ages of 40 and 60 tend to have higher systolic pressures than those between 20 and 40 it is not true to say that a normal blood pressure should be 100 mm Hg plus the age of the patient in years. On the contrary insurance companies well recognise the value of low figures and it is probable that the higher average pressures of the middle aged and elderly are due to atherosclerosis (Lewis 1938). The normal diastolic blood pressure lies between 60 and 90 mm Hg. The mean pressure approximates to the diastolic plus one third of the pulse pressure.

A common source of error in blood pressure estimation results from failure to obtain a basal reading this may be due to impatience or to lack of recognition of emotional or other physiological factors. Whenever the pressure is found to be raised the cuff should be left in position so that a second reading may be taken at the end of the examination. Casual measurements in healthy young adults who are a little anxious often register 160/90 mm Hg but if the patient is put at ease and allowed to rest quietly on a couch this figure may fall steadily to normal levels. It must be thoroughly understood that the maximum normal blood pressure of 145/90 mm Hg is basal. The question of pre hypertensive levels will be discussed later.

Slight disparity between readings taken from each arm is common especially in atherosclerotic and hypertensive subjects but the difference rarely exceeds 5 mm Hg (Amsterdam and Amsterdam 1943). The blood pressure is sometimes taken in the legs with the cuff above the knee and the stethoscope in the popliteal fossa. In the average normal individual in the horizontal position the blood pressure in the legs reads 20 to 40 mm Hg above that in the arms. This difference is lessened if the body is tilted head down and increased if it is tilted head up. The discrepancy appears to depend upon the cuff method of estimating the blood pressure for it is not found when the blood pressure is measured by means of direct arterial puncture (Loman *et al* 1936). In the standing position the systolic pressure in the arms measured at heart level usually shows no appreciable change but in 33 per cent of normal subjects it drops about 10 to 15 mm Hg the diastolic pressure rises about 5 mm Hg in 48 per cent of normal subjects drops about 5 mm Hg in 12 per cent and remains unchanged in 40 per cent (Currens 1948).

The ocular fundi Before leaving the peripheral vascular system the ocular fundi should be examined. The ophthalmoscope should be used with both eyes open and with either hand so that one may hold the instru-

ment with the right hand when examining the patient's right eye and with the left hand when examining his left eye. There are four features of particular interest to the cardiologist: the appearance of the disc, the calibre of the arteries, the presence of hæmorrhages and the presence of exudates. Details are described on page 423.

EXAMINATION OF THE HEART

Having gleaned as much information as possible from general inspection from searching for signs of failure and from examining the peripheral vascular system, one may turn with advantage to the heart itself and duly inspect, palpate, percuss and auscultate.

Inspection. The position and character of the cardiac impulse, if visible and of any other thoracic pulsation, should be noted. In this way, left or right ventricular hypertrophy, gallop rhythm, dilatation of the pulmonary artery and aortic aneurysm may be detected. Præcordial deformity may be observed and if due to the heart indicates its enlargement during the period of thoracic growth. Depression of the sternum or other thoracic deformity should be noted for it may alter the shape or position of the heart. Systolic indrawing of the thoracic wall is not abnormal if it occurs over the right ventricle and may be seen in the anterior axillary line when there is gross cardiac enlargement as a sign of adherent pericardium (see page 351); it should be looked for posteriorly over the last two ribs as described by Broadbent (1895).

Palpation. The apex beat, which is a geographical point, should be determined by locating the exact site of the maximum cardiac impulse. The physician's hand should be placed over the region of the fifth left intercostal space in the nipple line in order to ascertain its approximate position; the middle finger should then be directed vertically over it and shifted about until the maximum thrust is located. This rather than the lowest left point of such pulsation is the apex beat. Its position should be recorded with reference to the intercostal spaces, to the mid line and to the mid-clavicular line. It is usually in the fifth intercostal space, 8-9 cm. to the left of the mid line or just within the mid-clavicular line. If it is located beyond these confines, the possibility of displacement from scoliosis, elevation of the diaphragm or from pulmonary or pleural lesions should be considered before concluding that the heart is enlarged.

The character of the cardiac impulse is as important, if not more important than its position (the apex beat); it should be sensed both with the palm of the hand and with the finger tip. The qualities of heaving, thrusting, over action, tapping and triple rhythm can only be learned at the bedside.

Palpation may next be used to detect the presence of thrills, preferably in forced expiratory apnoea. This manœuvre brings the heart and great vessels closer to the chest wall, encourages the lung to retract from its buffering position and lessens the chance of confusing cardiac with respira-

tory phenomena. The vibration sense of normal individuals varies considerably, but increased perception comes with experience and good technique. Thrills should be timed against carotid pulsation.

Finally, palpation may be employed to detect abnormal pulsations of the great vessels, especially from aneurysm of the ascending aorta or from dilatation of the pulmonary artery.

Percussion. The value of percussing the heart has given rise to much dispute, many modern cardiologists maintaining that its place has been taken by the far more accurate and fertile method of radiography. The older school, however, modestly suggest that it is a useful bedside method, which gives reliable and helpful information if practised diligently, and if its limitations are appreciated. Certainly, if a fluoroscope is available, percussion is pointless, but a fluoroscope may not be available, or the patient may be so ill that only a portable X-ray machine can be used, and the distorted skiagram so obtained is liable to gross misinterpretation. In such cases percussion may be of value, and by constant practice the physician should learn what can and what cannot be expected from it.

The approximate position of the left border of the heart may be checked when the apex beat is difficult to locate, and dullness beyond the known or probable confines of the apex beat may sometimes be detected in cases of pericardial effusion.

It is impossible to determine the right border of the heart by percussion unless there is aneurysmal dilatation of the right auricle. On the other hand, pericardial effusion, even of moderate degree, can often be demonstrated.

It was once customary to speak of relative and absolute cardiac dullness, the latter being the note heard over the area of heart not covered by lung, but it is doubtful whether this distinction can be maintained. Diminution or absence of cardiac dullness, however, is a useful sign of emphysema.

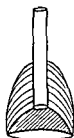
Percussion at the base may be rewarded in pericardial effusion, there being characteristic dullness in the second left interchondral space when the patient lies flat, also in substernal goitre, and in anterior aneurysm of the aorta, when a band of dullness extends laterally from the manubrium sterni.

Auscultation. When a man buys a tool for some specific purpose, he usually takes care that it is the best available for the particular job in hand. It is therefore strange that a superstition has grown up within the medical world that the older and more disreputable a stethoscope, the better, that it is not the stethoscope which matters, but the man behind it. This, of course, is nonsense. When a student fails to hear a murmur which is heard easily by another, exchange of stethoscopes quickly leads to mutual understanding. There is another curious tradition, fostered by many who appreciate the value of a good stethoscope, that the chest piece must be bell-shaped, and that any other type, especially the flat diaphragm (Bowles), is pernicious. This doctrine is as unreasonable as the first, for there is no

doubt that certain high pitched sounds, especially aortic diastolic murmurs and faint tubular breathing which can be heard with ease through a Bowles may be inaudible through a bell. The physical laws which govern auscultation have been studied by Rappaport and Sprague (1941). The diameter of the Bowles chest piece should be about $1\frac{1}{2}$ inches the cup should be shallow and its edge sharp (fig 101 a). Good material for the diaphragm is photographic or X ray film washed clean in hot water and cut to shape. The rubber tubing must be thick and should fit snugly to the connections. The internal calibre of the whole system should be nowhere less than the



(a) Bowles



(b) Bell

Fig 101—Binaural Stethoscopes

diameter of the hole in the centre of the chest piece which should measure about 5 mm. A good bell stethoscope (fig 101 b) is better for detecting low pitched sounds such as soft mitral diastolic murmurs moreover its range of sensitivity may be increased by varying the force with which it is applied to the chest wall. Light contact accentuates low pitched sounds firm pressure high pitched sounds. The cup should not be too deep and its diameter should not be less than one inch.

There are two other types of stethoscope which deserve comment the monaural wooden instrument of by gone days, and the differential stethoscope (symballophone). The rigid wooden stethoscope is rarely used nowadays but by combining

aural and tactile senses it facilitates the recognition of gallop rhythm. The differential stethoscope is constructed as shown in fig 102 and may be used to compare the timing of sounds originating at different sites and to determine the direction in which a murmur is propagated (Kerr *et al* 1937).

Auscultation of the heart can only be learned at the bedside but the following advice may be helpful to students. The præcordium should be examined all over not just at areas where individual valve sounds are expected gallop rhythm pericardial friction and certain important murmurs will not then escape notice. It is enough to listen to one thing at a time thus when an expert hears a soft elusive mitral diastolic murmur hitherto overlooked it is not necessarily because he has better ears or a better stethoscope but because he has acquired a more selective power of concentration. Basal murmurs and pericardial friction are heard most easily in expiratory apnoea tricuspid murmurs in inspiratory apnoea mitral murmurs in the left lateral position especially after exercise or after the inhalation of amyl nitrite. Heart sounds should be timed against carotid pulsation if difficulty is experienced due to tachycardia the heart may be slowed by carotid sinus compression.

The heart valves lie so close together that a stethoscope placed over them transmits sounds from all making it difficult to distinguish one from another. Certain favourable areas have come to be recognised however at which each valve may be studied selectively. Although such areas represent maximum purity of particular valve sounds they do not represent maximum intensity. Thus if a valve sound or murmur is faint it may be inaudible at the site of selection yet heard clearly elsewhere. Aortic diastolic murmurs for example with the possible exception of those associated with

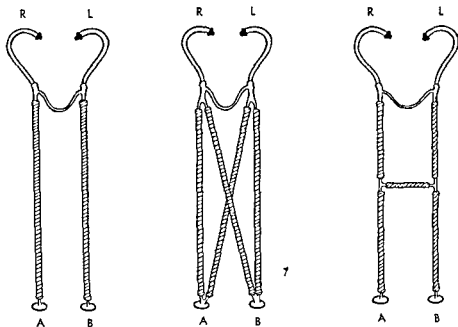


Fig. 102—The Differential Stethoscope (3 varieties). Sounds travelling from A to B reach the right ear before the left giving the impression of movement in that direction.

syphilitic aortitis are usually heard much better down the left border of the sternum than at the aortic area (second right costal cartilage). Again both aortic systolic and diastolic murmurs are occasionally maximal at the mitral area (apex beat). It follows that analysis of cardiac sounds and murmurs requires more than geographical data. Whilst the propagation of a basal murmur may help to distinguish between aortic and pulmonary responsibility, propagation of a central or apical murmur bears more relation to the intensity of the sound than to its site of origin and is of little value in differential diagnosis. The quality and timing of murmurs however are very important and will be discussed later.

The Heart Sounds The first heart sound is due almost entirely to mitral and tricuspid valve closure. Its intensity depends chiefly upon the position of the cusps at the beginning of ventricular systole partly upon the volume

of lung covering the heart or upon the thickness of the chest wall and perhaps least upon the degree of ventricular filling and the strength of ventricular contraction — unless changes in these functions are extreme

The position of the valve leaflets at the beginning of ventricular systole depends upon the P R interval the loudest first sound is produced when auricular contraction forces the leaflets wide open immediately before the ventricles contract (P R around 0.10 second) but when there is a relatively long delay between auricular and ventricular excitations ($P-R=0.2$ second) the cusps may float into apposition before the ventricles contract and hence shut quietly (Dock 1933) The loud first sound of the hyperkinetic circulatory states is similarly produced in that a relatively high and effective filling pressure keeps the cusps wide open until the last possible moment whatever the auricles are doing

The second heart sound is due to aortic and pulmonary valve closure Although the aortic element may be heard best in the aortic area in the neck and at the apex beat and the pulmonary element in the pulmonary area no such distinction is entirely reliable In normal individuals both elements can usually be heard especially in the second and third left interchondral spaces close to the sternal border In children and adolescents the split is often obvious (grade II) particularly towards the end of inspiration The first element is aortic the second pulmonary In adults recognition of splitting may be more difficult (grade I) but becomes easier with experience Pathological splitting (grade III) is due to delay in pulmonary valve closure and is usually due to right bundle branch block or possibly to delay in the emptying time of an over filled right ventricle Slight delay in aortic valve closure may bring the two elements together and so cause a single second heart sound There seems to be a tendency for this to occur with advancing years With left bundle branch block the aortic element may lag behind the pulmonary element but rarely so much as to cause more than reversed grade I splitting

Recognition of a split second heart sound at once proves that both semilunar valves are functioning and thus excludes pulmonary stenosis Recognition of grade III splitting is also helpful because the knowledge that right bundle branch block is probably present may be of considerable diagnostic importance

Accentuation of the second heart sound may result from systemic or pulmonary hypertension the clinical circumstances rather than the site of maximal intensity usually decide which although it may be possible to tell directly if there is also wide splitting If the ascending aorta or pulmonary artery is unusually close to the anterior surface of the chest either because it is abnormal or because it is scantily covered by lung and chest wall the second heart sound is also loud Conversely a soft or absent second heart sound is usual in emphysema

The third heart sound and other varieties of triple rhythm are discussed elsewhere (page 164)

Examination of the lungs Whilst routine examination of all systems is necessary in any speciality examination of the lungs is peculiarly significant in cardiology for several reasons to detect any abnormality associated with the pulmonary circulation e.g. congestion œdema infarction hydrothorax and the like to see if there is any pulmonary cause for anoxæmia especially emphysema to note whether there are any changes in the lungs which might be secondary to bronchial obstruction due to pressure from some cardiovascular lesion such as pericardial effusion or aneurysm of the aorta to reveal any thoracic or intra thoracic cause for cardiac displacement e.g. scoliosis pulmonary collapse or fibrosis pleural effusion pneumothorax or elevation of the diaphragm

Examination of the other systems There is no need to go into these in detail but it must be recognised that important clues to cardiovascular diagnosis lie outside that system Thus atrial septal defect may be suggested by arachnodactyly rheumatic activity by erythema marginatum diphtheritic carditis by paresis of accommodation or palatal palsy beri beri heart and periarteritis nodosa by polyneuritis bacterial endocarditis by the combination of fever anæmia petechiæ splenomegaly and clubbing of the fingers thyrotoxic heart disease by remote findings in the eyes neck and hands syphilitic aortitis by signs of syphilis elsewhere especially in the central nervous system and so forth

SPECIAL TESTS

History taking and routine physical examination are supported by fluoroscopy and electrocardiography These two methods of investigation are so important that a special chapter is devoted to each (Chapters II and III) There are however certain other tests which may be used with advantage in appropriate circumstances and which may be described conveniently here some are beyond the scope of the general practitioner but are included to foster interest and understanding

Direct measurement of the venous blood pressure Whilst elevation of the venous pressure is usually detected clinically with little difficulty there are occasions when it is valuable to check it by direct measurement The subject should be propped up at an angle of 45 degrees because patients with orthopnoea cannot lie flat and the technique should be the same for all cases The right arm bared to the shoulder is abducted to a right angle and supported on pillows so that the antecubital fossa is roughly at heart level An infusion needle connected to a spinal manometer or similar graduated glass tube is then inserted into the antecubital vein the zero mark on the manometer being placed at the level of the fourth costal cartilage by means of a spirit level the height to which blood rises above this mark represents the venous pressure Alternatively the zero mark may be placed at the level of the sternal angle or of some other reference point To avoid clotting a saline reservoir containing a drop of heparin should be

rapidly through a wide bore needle the patient having been warned to raise the other hand smartly the instant he should notice a strange taste in or under the tongue. This taste is peculiarly intense and bitter so that it is difficult for the patient to be mistaken about the moment of its arrival and objective confirmation may be obtained by the involuntary grimace which accompanies it. The time should be measured from the beginning of the injection to the end point described and in normal individuals averages 13.5 seconds with extremes of 9 to 18 seconds (Wood 1936). A concentrated solution of saccharin (2.5 G in 4 ml of water) 5 ml of 10 per cent calcium gluconate and numerous other substances may be used instead of decholin but none have a more definite end point. Saccharin is less unpleasant however does not cause vomiting and may be injected repeatedly when serial observations are required.

The arm to lung time may be measured by injecting 0.25 ml of ether into the antecubital vein its arrival in the capillaries of the lung being signalled by a sudden cough or deep breath and by the smell of ether in the expired air. Amyl acetate may also be used the smell of pear drops being unmistakable when it reaches the lungs. The normal time averages 6 seconds and ranges between 3.5 and 8 seconds (Hitzig 1935). The test has a limited value as explained on page 163.

The passage of radioactive sodium through the chambers of the heart may be recorded in the form of a time concentration curve by means of a Geiger Muller counter specially fitted with a direct writing attachment (Prinzmetal *et al* 1948).

CARDIAC CATHETERISATION

Although first performed by Forssmann (1929) on himself the introduction of cardiac catheterisation as an aid to clinical diagnosis is largely due to the work of Cournand in the U.S.A. and of McMichael and Sharpey Schafer in England. The median cubital vein of either arm is exposed and the tip of a special nylon catheter inserted in the manner of introducing a cannula. The catheter which has been previously washed through with saline and in the hilt of which is an adaptor connected with a 5 or 10 ml syringe loaded with saline is then pushed up the vein under fluoroscopic control its curved tip being directed medially. Any obstruction may be overcome by rotating the catheter a little one way or the other. Hooks at the thoracic inlet may be passed during deep inspiration sometimes it may be necessary to exchange the catheter for one with a more curved tip or even to use the other arm. Venospasm is avoided by proper skin anaesthesia and by choosing a catheter that is not too large for the vein. When the tip of the catheter lies in the right auricle the syringe is removed and the adaptor is connected by means of a rubber tube and Y piece to a saline manometer and reservoir into which a drop of heparin is added. The catheter is flushed from time to time by releasing the clip below the reservoir or a continuous drip technique may be employed. The



Fig 1 04 (top)—Catheter in right branch of the pulmonary artery

Fig 1 05 (bottom)—Catheter in right branch of the pulmonary artery (1st oblique position)



Fig 1 06—Catheter in right branch of the pulmonary artery (2nd oblique position)



Fig 1 07—Catheter in left branch of the pulmonary artery



Fig 1 08—Catheter in left branch of the pulmonary artery (1st oblique position)

best way to enter the right ventricle is to loop the catheter in the right auricle and to rotate the upwardly directed tip until it faces the tricuspid orifice. If it can then be passed through the valve it is in the right position for entering the pulmonary artery. Success is signalled by a sudden rise of pressure in the manometer and by increased amplitude of pulsation. The catheter is then advanced into the pulmonary artery (figs 104-109). With a saline manometer only mean pressures can be recorded. Electrical manometers with optical recording overcome this difficulty and provide continuous graphic records showing systolic and diastolic levels (Hamilton *et al* 1934). Ten ml samples of blood may be taken under paraffin from appropriate chambers for estimation of oxygen unsaturation.

Cardiac catheterisation is of value as a means of obtaining samples of mixed venous blood for estimating the cardiac output in demonstrating certain shunts and in measuring the pressures in the right side of the heart. If 5 litres of blood pass through the lungs per minute and the arteriovenous oxygen difference is 50 ml per litre then the amount of oxygen taken up by the lungs is $5 \times 50 = 250$ ml per minute. In other words the cardiac output in litres per minute \times the arterio venous oxygen difference in ml per litre = the oxygen consumption in ml per minute. This principle was first enunciated by Fick (1870) and is usually expressed as follows

$$\text{Cardiac output (litres per min)} = \frac{\text{Oxygen consumption (ml per min)}}{\text{Arterio venous oxygen difference (ml per litre)}}$$

The oxygen consumption may be measured by means of a spirometer in the usual way. The arterio venous oxygen difference is estimated by measuring the oxygen unsaturation of 5 or 10 ml samples of blood from the right auricle and from the femoral artery obtained by means of cardiac catheterisation and direct arterial puncture respectively subtracting one



Fig 109—Catheter in left branch of the pulmonary artery (2nd oblique position)

from the other and expressing the result in ml per litre. Blood gas analysis requires familiarity with Haldane's or van Slyke's apparatus. The normal resting cardiac output is about 5 litres per minute in the horizontal position and 4 litres per minute in the erect position (McMichael and Sharpey Schafer 1944). During exercise it may rise to 20 to 30 litres per minute.

Intra cardiac shunting may be proved by finding significant differences in the degree of oxygen unsaturation in samples of blood taken from the *venæ cavæ* and right auricle as in atrial septal defect or from the right ventricle and pulmonary artery as in patent ductus. In a normal subject with 100 per cent hæmoglobin the arterial blood contains about 190 ml oxygen per litre as the maximum oxygen capacity is about 200 ml per litre (page 17) this gives 10 ml oxygen unsaturation per litre which is the figure obtained by gas analysis. Mixed venous blood usually gives an unsaturation value of about 60 ml per litre this may be expressed as $200 \text{ minus } 60 = 140 \text{ ml per litre oxygen content}$ or as

$\frac{140}{200} \times 100 = 70 \text{ per cent oxygen saturation}$. Complete mixing is always found in the pulmonary artery and usually in the right ventricle but stream lining in the right auricle may give rise to false readings if there is an appreciable difference between superior and inferior *vena cava* samples. Proof of an intra cardiac shunt is rarely accepted unless the oxygen unsaturation of samples from different parts of the right side of the heart differ consistently by more than 10 ml/litre.

The normal mean right auricular pressure lies between -2 and 2 mm of mercury with reference to the centre of the auricle in the recumbent position or between -8 and -4 cm of saline with reference to the sternal angle. The effective filling pressure of the heart is the auricular pressure minus the negative intra thoracic pressure i.e. $0 - (-5) \text{ mm Hg} = +5 \text{ mm of mercury}$. The mean right ventricular pressure is 10 to 15 cm of saline above the right auricular pressure the right ventricular systolic pressure is 20 to 30 mm of mercury (Bloomfield *et al* 1946). The mean pressure in the pulmonary artery is about the same as that in the right ventricle the pulmonary diastolic pressure being very low. Measurement of such pressures may provide valuable information in diagnosis and research.

The chief risks of cardiac catheterisation are paradoxical embolism in cases of veno arterial shunt rigors due to pyrogens in the apparatus and subsequent thrombo phlebitis. Paradoxical embolism may cause hemiplegia or cerebral abscess and great care must be taken when investigating cases of right to left shunt heparin should be given through the catheter at the start in a dose of 25 to 50 mg according to the age and weight of the patient ($1 \text{ mg per kg body weight}$) and air must be rigidly excluded from the system especially when changing syringes during sampling. Rigors can be very dangerous in cases of severe mitral stenosis or left ventricular failure when the resulting rise of venous pressure may precipitate an attack.

of acute pulmonary œdema. As the rubber connecting tubes carry saline only, the pyrogens may be assumed to come from the lining of the catheter which should either be destroyed after a rigor or thoroughly cleaned with hydrogen peroxide. Thrombo phlebitis is rarely serious but may give rise to transient swelling of the arm and occasionally to pulmonary infarction. Infection may be avoided by careful aseptic technique and by giving a massive dose of penicillin immediately after the procedure. In cases with right to left shunt thrombo phlebitis should be treated with heparin without delay (see page 454).

Arterial oxygen saturation. With a normal hæmoglobin of 15 gm per cent one litre of blood should contain $15 \times 1.34 \times 10 = 201$ ml of oxygen when fully saturated because 1 G of hæmoglobin is capable of holding 1.34 ml of oxygen. As normal arterial blood is 95 per cent saturated it contains at N.T.P. roughly 190 ml of oxygen per litre neglecting a small amount held in solution in the serum. The demonstration of an abnormal degree of arterial oxygen unsaturation is helpful in proving the existence of a veno arterial shunt or of some disease of the lungs causing failure of proper oxygenation of the blood in the pulmonary circulation e.g. severe emphysema.

Samples of arterial blood may be obtained from the femoral artery by direct puncture. An ordinary intramuscular needle attached to an all glass syringe containing a little paraffin should be plunged almost vertically into the vessel. The femoral blood pressure then lifts the barrel of the syringe which fills with blood accordingly no aspiration is necessary. On withdrawing the needle the site of puncture should be compressed digitally for at least a minute. Some risk is attached to the procedure if the vessel is unduly sclerotic when a severe hæmatoma may result. The sample of arterial blood is analysed as mentioned in the preceding section.

Vital capacity. This is easily measured by means of a spirometer. During normal respiration the amplitude of the graph represents the tidal air usually about 500 ml. The complemental air is registered by the additional upright deflection inscribed when the subject takes a maximum inspiration. The reserve air is represented by the additional downward deflection inscribed when the subject takes a maximum expiration. Each averages about 1.500 ml. The vital capacity is the sum of these three measurements (fig. 1.10) and averages 3½ to 4 litres being higher in men than in women. It has been found to be proportional to the surface area of the body which may be calculated from height and weight by reference to standard tables. The vital capacity is decreased particularly in emphysema and in heart failure with pulmonary congestion.

Lung volume and residual air. The residual air is that which remains in the lungs after a complete expiration and may be estimated in the following way. The patient is connected with a spirometer containing known quantities of air and oxygen and encouraged to breathe quietly and to become accustomed to the instrument. The vital capacity and reserve air are

CC

0

500

1000

1500

2000

2500

3000

3500

4000

4500

1 Hand

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

R

Y

X

C

V

measured in the usual way. Oxygen is admitted at a rate equivalent to its uptake so that the graph remains flat. The patient is then disconnected and a known quantity of helium (hydrogen is not altogether safe) is introduced into the system. The percentage of this gas in the spirometer is next measured by means of a katharometer (McMichael 1939) and should tally with the calculated percentage. When the patient is again connected to the circuit helium diffuses into the lungs until equilibrium is reached. The capacity of the space into which the gas has diffused may be calculated from the fall in the percentage of helium in the spirometer. With certain corrections this represents the reserve air plus the residual air. As the former is already known the latter may be calculated by subtraction. The total lung volume is the vital capacity plus the residual air.

In emphysema reduction of vital capacity is balanced by an increase in the residual air so that the total lung volume remains normal. In pulmonary congestion however the vital capacity is reduced at the expense of the total lung volume, the residual air remaining unchanged and is a measure of the amount of extra blood held in the pulmonary circulation.

The jugular phlebogram. The polygraph is an instrument for making simultaneous graphic records of two or more vascular pulsations. MacKenzie (1902) used the ink polygraph and concentrated on the jugular

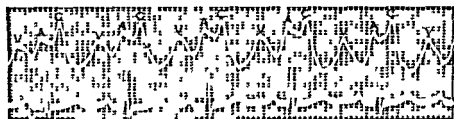


Fig. 111—Jugular phlebogram

A A 1 co ct
C V t cul on t d losur f th p d l
V Th nm t gnal th p g of th i p d l

phlebogram as a means of analysing abnormalities of rhythm. Although the instrument is now rarely used in clinical work owing to the development of the more fruitful electrocardiograph it is still necessary to understand the significance of the venous waves in the neck. There are three in each cardiac cycle labelled a, c and v (fig. 111). The first represents auricular contraction and disappears when the auricles fibrillate. It may be distinguished clearly in many cases of heart block as an isolated event between two main jugular beats. The c wave signals isometric ventricular contraction and closure of the tricuspid valve. In the neck it merges into the artefact caused by carotid pulsation. The v wave occurs after a longer interval; its rising limb signifies increasing venous pressure resulting from continued venous return against a closed tricuspid valve and reaches its peak the instant before the valve opens when the pressure promptly falls. The

summit of the v wave is therefore used as a means of timing the opening of the tricuspid valve, and therefore of the mitral. The reduction of venous pressure causing the trough (x) after the a wave is due to auricular relaxation. The trough (x_1) between c and v represents the negative pressure created by descent of the atrio-ventricular septum and marks the systolic ejection phase. The final trough (y) following the summit v is due to release of auricular pressure following the opening of the tricuspid valve.

Simultaneous jugular phlebograms, electrocardiograms and phonocardiograms have helped considerably in the elucidation of gallop rhythm of the third heart sound and of the opening snap of mitral stenosis. Optical recording has replaced the graphic method employed by Mackenzie so that errors due to the inertia of levers is minimised.

The arteriogram. There are numerous methods of recording pulsation from any superficial artery but those in which the graph is optically registered are preferable. They are of limited clinical value because they reveal little which cannot be discerned with the trained finger. A normal arteriogram (fig. 112) usually exhibits two waves P and D . The former is the percussion wave and represents the rapidly transmitted shock of left

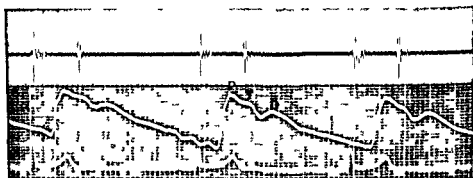


Fig. 112—Brachial arteriogram

P P cu D D rot w T T d w e

(B) ur ty f Dr F G d e r a d M Zoob)

ventricular contraction it is a pressure wave and must not be confused with blood flow. Its velocity is 5 to 8 metres per second and is inversely proportional to the elasticity of the artery. The length of the wave is 3.5 to 5 metres. D is the dicrotic wave and is produced by the shock of aortic valve closure. The latter synchronises with the incisura N (dicrotic or aortic notch) which precedes the dicrotic wave. Under certain circumstances e.g. in combined aortic stenosis and incompetence a second systolic wave T follows the percussion wave. This is believed to be a tidal wave the tail of the percussion wave being suddenly augmented by the reflection of its head from the periphery (Bramwell 1937).

Phonocardiography. The heart sounds were first directly recorded by Frank (1904) and modifications of his original methods are still widely

used. A stethoscope is applied to the chest wall and connected through a rubber tube with a Frank's capsule. This consists essentially of a very thin rubber membrane to which a small mirror is attached. A beam of light is focused on the mirror and reflected onto a camera unit. Electrical amplification was developed by Einthoven (1907) and has been employed in many ways since but has little advantage over the direct method (Orias and Braun Menendez 1939). Tracings should be timed against the jugular phlebogram which is the simplest means of recording all mechanical events in the cardiac cycle (fig. 113).

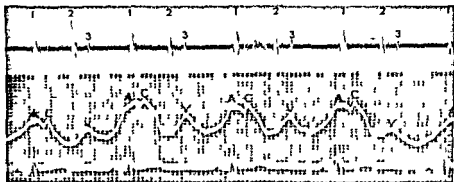


Fig. 113—Phonocardiogram synchronised with electrocardiogram and jugular phlebogram. Note the third heart sound and absence of murmur.

(By courtesy of Dr. Francis Gardner and Mr. Zook.)

Phonocardiography has been largely restricted to academic and research units but deserves to be employed more widely as a routine aid to diagnosis. It has proved valuable in the elucidation of extra heart sounds and in the precise timing of murmurs.

Ballistocardiography. The ejection of blood from the heart is accompanied by a recoil of the whole body in the opposite direction proportional to the mass ejected. If the body is placed horizontally on a specially constructed couch the recoil movements may be graphically recorded. The ballistocardiogram has been used chiefly to measure changes in cardiac output but the form of the graph may give other information about the function of the heart (Starr 1941; Starr and Mayock 1948).

GENERAL TESTS

Of more general tests, examination of the urine, renal function tests, the blood count, the blood Wassermann reaction and the Kahn reaction, and the basal metabolic rate are most valuable in cardiovascular work. Their technique will not be described but it is pertinent to make a few remarks concerning them.

Examination of the urine. The urinary findings may be helpful in cardiology in four main ways. First, albuminuria is commonly present in con-

gestive heart failure the diagnosis of which should be carefully checked in its absence. Second diabetes mellitus may be associated with coronary or peripheral atherosclerosis. Third red cells are nearly always found in the urine during the course of bacterial endocarditis and their repeated absence is enough to question the diagnosis the positive finding is less significant for red cells may appear in the urine in many other conditions. Fourth the urinary findings in hypertension are of paramount importance.

Renal function tests The blood urea, urea clearance and urine concentration tests are those commonly employed. Their chief value is in hypertensive heart disease. It should be borne in mind that the blood urea is often raised up to about double the normal in congestive heart failure from any cause with a corresponding fall in urea clearance. Renal function tests are usually normal in embolic nephritis and in renal infarcts.

Blood count Increased activity of the red marrow with reticulocytosis is frequent in congestive heart failure especially when associated anoxæmia can be demonstrated as in pulmonary heart disease. Polycythæmia is far less common except in congenital heart disease with cyanosis. The effect of anæmia on the heart is described in Chapter XX. Here it is only necessary to say that the anæmia which may be associated with rheumatic or bacterial endocarditis and with malignant or chronic nephritic hypertension is usually normocytic and orthochromic. Anæmia associated with myxædema is variable and may be orthochromic or hypochromic in the latter however iron deficiency is usually present as well.

Leucocytosis may occur in rheumatic carditis and in bacterial endocarditis but normal white counts are common in both and indeed leucopenia occurs in about one third of cases of bacterial endocarditis. The leucocytosis of myocardial infarction is transient lasting little longer than the fever. Repeated white counts are necessary as a check against the development of agranulocytosis during the course of thiouracil therapy and during prolonged prophylactic treatment with sulphonamides.

The erythrocyte sedimentation rate The E.S.R. is helpful in cardiology in several ways it is the most sensitive guide to the presence of activity in rheumatic carditis it is accelerated in syphilitic aortic incompetence which may thus be distinguished from old rheumatic and atherosclerotic varieties its characteristic behaviour in myocardial infarction serves as a useful index of progress its acceleration in the various forms of hypertension runs more or less parallel to the degree of renal involvement being rapid in nephritic and malignant hypertension and normal in the benign type it is retarded by congestive heart failure the development of which may suddenly alter a reading of 80 or so in rheumatic carditis for example to one below 10. Readings are normal in congenital heart disease (or low if there is polycythæmia) in old rheumatic heart disease in uncomplicated angina pectoris in thyrotoxicosis and in myxædema (Wood 1936).

The basal metabolic rate As thyrotoxicosis and myxædema are not uncommon causes of cardiovascular disorder and thiouracil is sometimes

used to induce myxœdema in stubborn cases of angina pectoris and congestive heart failure the B M R is frequently measured in cardiovascular clinics. It is important therefore to understand that readings are often elevated in heart failure from any cause if there is dyspnœa for under such circumstances truly basal conditions cannot be realised.

REFERENCES

- Amsterdam B and Amsterdam A L (1943) Disparity in blood pressures in both arms in normals and hypertensives and its clinical significance: a study of 1 000 normals and 272 hypertensives. *N Y State J Med* 43 2294
- Bloomfield R A, Lanson H D, Cournand A, Breed E S and Richards D W (1946) Recording of right heart pressures in normal subjects and in patients with chronic pulmonary disease and various types of cardiocirculatory disease. *J clin Invest* 25 639
- Bramwell C (1937) The arterial pulse in health and disease. *Lancet* ii 39
- Broadbent W (1895) An unpublished physical sign. *Ibid* ii 200
- Currens J H (1948) A comparison of the blood pressure in the lying and standing positions: a study of five hundred men and five hundred women. *Amer Heart J* 35 646
- Dock W (1933) Mode of production of the first heart sound. *Arch Int Med* 51 737
- Einthoven W (1907) Die Registrierung der menschlichen Herztöne mittel des Saiten galvanometers. *Pfugers Arch ges Physiol* 117 461
- Fick A (1870) Ueber die Messung des Blutquantums in den Herzventrikeln. Sitzungsberichte der phys. med. Gesellsch. zu Würzburg.
- Förssmann W (1909) Die Sondierung des rechten Herzens. *Klin Wchnschr* 8 2085
- Frank O (1904) Die unmittelbare Registrierung der Herztöne. *Munch med Wchnschr* 51 953
- Hamilton W F, Brewer G and Brotman I (1934) Pressure pulse contours in the intact animal. I—Analytical description of a new high frequency hypodermic manometer with illustrative curves of simultaneous arterial and intracardiac pressures. *Amer J Physiol* 107 427
- Hitzig W M (1935) The use of ether in measuring the circulation time from the antecubital veins to the pulmonary capillaries. *Amer Heart J* 10 1080
- Joint Report of the Committees Appointed by the Cardiac Society of Great Britain and Ireland and the American Heart Association (1939) Standardization of methods of measuring the arterial blood pressure. *Brit Heart J* 1 61
- Kerr W T, Althausen T L, Bassett A M and Goldman M J (1937) The symbolophone: a modified stethoscope for the lateralisation and comparison of sounds. *Amer Heart J* 14 594
- Lewis W H Jr (1938) Changes in age in the blood pressure of adult men. *J Physiol* 1-2 401
- Loman J, Dameshek W, Myerson A and Goldman D (1936) Effect of alterations in posture on the intra arterial blood pressure in man. I. Pressure in the carotid, brachial and femoral arteries in normal subjects. *Arch Neurol Psychiat* 35 1-16

Mackenzie J (1902) The study of the pulse Edinburgh

McMichael J (1939) A rapid method of determining the lung capacity
Clin Sc 4 167 — Sharpey Schafer E P (1944) Cardiac output in man by
a direct Fick method *Brit Heart J* 6 33

Orias O and Braun Menendez E (1939) The heart sounds in normal and
pathological conditions London

Prinzmetal M Corday E Bergman H C Schwartz L and Spritzler R J
(1948) Radiocardiography A new method for studying the blood flow through
the chamber of the heart *Science* 108 340

Rappaport M B and Sprague H B (1941) Physiologic and physical laws
that govern auscultation and their clinical application *Amer Heart J* 21 257

Robb G P and Weiss S (1934) The velocity of pulmonary and peripheral
venous blood flow and related aspects of the circulation in cardiovascular disease
Ibid 9 742

Starr I (1941) Clinical studies with the ballistocardiograph incongestive
failure on digitalis action on changes in ballistic form and in certain acute experi-
ments *Amer J med Sci* 202 469 — Mayoock R L (1948) On the signi-
ficance of abnormal forms of the ballistocardiogram A study of 234 cases with 40
necropsies *Ibid* 215 631

Weiss S Robb G P and Blumgart H L (1928) The velocity of blood
flow in health and disease as measured by the effect of histamine on the minute
vessels *Amer Heart J* 4 664

Winternitz M Deutsch J and Brull Z (1931) Eine klinische brauchbare
Bestimmungsmethode der Blutemlaufzeit mittels Decholinjektion *Med Klin*
27 986

Wood P H (1936) The erythrocyte sedimentation rate in diseases of the
heart *Quart J Med* 29 1 — (1936) Right and left ventricular failure a
study of circulation time and venous blood pressure *Lancet* ii 15

CHAPTER II

RADIOGRAPHIC DIAGNOSIS

TECHNIQUE

THERE are at present five radiological methods applicable to cardiology fluoroscopy orthodiagraphy teloradiography kymography and angiocardiology Fluoroscopy (screening) is a routine diagnostic procedure orthodiagraphy is the construction of a simple tracing of the size and shape of the heart in any specified position as a supplement to fluoroscopy teloradiography gives a more accurate record and should be preferred when facilities permit kymography is a method of recording the character and amplitude of cardiac pulsation angiocardiology is the study of individual cardiac chambers or vessels with the aid of intravascular contrast media

FLUOROSCOPY

With modern X ray equipment a remarkably clear view of the heart may be obtained The patient should be stripped to the waist and pressed close to the viewing screen The diaphragm which controls the diameter of the beam emitted from the X ray tube is first opened wide in order to view the thoracic contents as a whole In this preliminary survey attention is paid to the lungs to the costo phrenic angles and to the general size and shape of the heart The diaphragm is then constricted so that only the heart can be seen and the latter is observed more critically The size shape and pulsation of each part should be noted in regular sequence On the right side (fig 201) a faint slightly concave line representing the superior vena cava descends from the sterno clavicular region close to the shadow of the vertebral column until it meets the ascending aorta which both displaces it to the right and causes it to become convex Below is the border of the right auricle which usually meets the diaphragm at a slightly acute

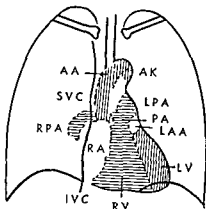


Fig 201—Diagram of antero poste or view of the heart as seen fluoroscopically

AA	Asc d g rt
PA	Pulm ry rt ry
RV	Rght rt l
RA	Rgh rt l
AK	A t k kl
LV	L f t l
LAA	L ft u cl pp d g
IVC	I f

angle The left border of the normal heart is made up of three convex curves from above downwards these are the aortic knob or knuckle, the pulmonary arc and the contour of the left ventricle Between the last two there is a small neutral segment or point of opposing movement which marks the left auricular appendage above it the pulmonary artery expands during systole while below the left ventricle contracts The hilar shadows are chiefly vascular the right pulmonary artery may be seen dividing early into upper and lower branches the former being indistinct the latter sweeping downwards in a well defined arc the left limb of the pulmonary artery forms the main pulmonary arc described above

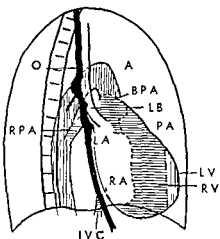


Fig 20—Diagram of right anterior oblique view of the heart as seen fluoroscopically (1st oblique position)

O	B	um	hil	d	ph	gu
BP A	Btu	t	n	f	th	p
LA	L	ft	ry	l		
A	A	t	rch			
I B	I	ft	bron	h		
IVC	I	f				
V	V	nt	l	s	(L	d R)
PA	Pulm	ry	rc			
RA	R	ght	a	r	l	

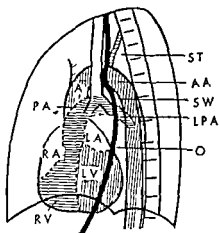


Fig 203—Diagram of left anterior oblique view of the heart as seen fluoroscopically (2nd oblique position)

RV	R	ght	ntr	l
LA	L	ft	nt	l
AA	A	rt	a	h
SW	S	b	ort	w
O	O	ar	um	fill
RA	R	ght	ur	l
LA	L	ft	ur	l
PA	P	ulmo	ry	h
ST	S	p	t	ngl

The patient is then turned into the first or right anterior oblique position. The observer should place his gloved hands on the patient's hips and manually rotate him (so that the right shoulder is brought to the front) until the position is satisfactory. The arms should be extended the left forwards and outwards the right backwards and outwards. In this view (fig 20-) the ventricular shadows are superimposed and the right auricle is rotated towards the front so that little can be learned about these three chambers on the other hand the left auricle is outlined clearly as it forms the upper part of the posterior border of the heart. Just anterior to the top of the left auricular curve a rather dense round shadow may be seen due to the bifurcation of the pulmonary artery it is connected with the anterior

ventricular border by a convex line representing the root of the pulmonary artery and conus of the right ventricle. Above it are the superimposed shadows of the ascending and descending parts of the aortic arch. If the patient is made to swallow a barium emulsion of the consistency of thick cream the œsophagus is outlined at the back of the heart. Under favourable conditions it is indented in turn by the arch of the aorta, by the pulmonary artery and left bronchus, and by the left auricle. The left bronchus may be seen between the œsophagus and the rounded shadow of the dividing pulmonary artery. Between the œsophagus and the vertebral column there should be a translucent space.

In the second or left anterior oblique position (fig 2 03) the patient is turned to the right through an angle of about 45 degrees, the left shoulder being brought forwards. In this view the two ventricles appear side by side, the left forming the posterior border of the heart shadow and the right the anterior, so that their contours can be readily compared. The shadow of the right auricle overlaps that of the right ventricle, the shadow of the left auricle lies posteriorly above the left ventricle. Cranially the aorta and pulmonary artery may be seen as two arches, one above the other, separated by a light space known as the sub aortic window, and crossed by the translucent trachea and left bronchus. The aortic arch and descending aorta are well defined and shaped like an inverted J, but the pulmonary artery is less distinct. Above the aorta is another light space, the supra aortic triangle, bounded by the vertebral column posteriorly, by the left subclavian artery anteriorly, and by the aortic arch below. The barium filled œsophagus is deflected to the patient's right as it crosses the aortic arch, then lies in close relation to a short segment of the descending aorta, leaves that vessel at about the level of the pulmonary artery, and courses downwards and to the subject's right across the shadow of the left ventricle.

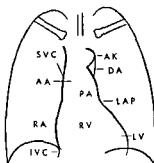


Fig 2 04—Orthodiagram of a normal subject (A.P. view)

ORTHODIAGRAMM

Clips should be fitted to the viewing screen to enable tracing paper to be held firmly in position. To make an accurate tracing of the heart shadow or orthodiagram (fig 2 04) special attention should be paid to five points. First the position of the patient must be properly adjusted to the view required; he must be pressed firmly against the screen and he should hold on to some support so that he can remain still. Second the tracing should be made in mid inspiration and as it cannot be completed in one period of breath holding, lines which move with respiration should be checked.

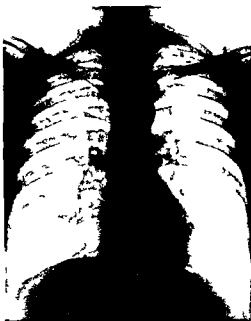


Fig 2 05—Teleradiogram of a normal subject (A P view)



Fig 2 06—Teleradiogram of a normal subject (1st oblique position)



Fig 2 07—Teleradiogram of a normal subject (2nd oblique position)

more than once. Third to avoid distortion the cardiac outline must be traced by means of parallel rays this is accomplished by constricting the diaphragm to the smallest aperture consistent with adequate visualisation. Fourth the greatest accuracy must be maintained when tracing the interior thoracic wall at its widest point and the lateral borders of the cardiac shadow, so that the cardio thoracic ratio is reliable. Fifth the finished orthodiagram should be checked against the shadows traced to make sure the patient has not moved during the procedure.

Fluoroscopes suitable for cardioscopy are so constructed that the X ray tube may be moved easily in any direction by the lever which operates the diaphragm. In making the tracing the small light spot is run swiftly over the contours of the heart great vessels clavicles interior thoracic wall and diaphragm. With experience it may be completed very quickly without danger of over exposing the patient or over heating the tube nevertheless a good technician switches off the current whenever momentarily disengaged. Fluoroscopy and orthodiagraphy are usually carried out with a power of 60 kilovolts and a current of 3 to 4 milliamps but with obese subjects it may be necessary to step up the kilovolts to 65 or 70 in order to obtain sufficient penetration. Tracings are made with a wax pencil and it is helpful to add signs denoting the degree and direction of pulsation of important chambers and vessels.

TELERADIOGRAPHY

Skiagrams of the heart are always taken at a tube screen distance of at least 6 feet preferably of 7 feet to avoid distortion by diverging rays. The duration of exposure should be half a second to ensure a diastolic record. To obtain a systolic record a very short exposure and a special timing device are required. Skiagrams of the oblique views are best taken when the most informative degree of rotation has been ascertained by previous fluoroscopy. The normal appearances are illustrated in figures 2.05-2.07.

KYMOGRAPHY

A specially constructed kymograph may be attached to a teleradiograph for the purpose of recording cardiac pulsation (Stumpf 1931). A lead screen containing horizontal slits 11 mm apart is interposed between the film and the patient's chest and made to descend 1 cm during one complete cardiac cycle. The timing of the exposure is adjusted to synchronise with the descent of the grid. In kymograms so obtained the lateral borders of the heart and great vessels appear toothed like the edge of a saw (fig 2.08) the ventricular crests representing diastole the troughs systole. Pulsation is recorded in only one dimension i.e. in a plane parallel to the film but if the three standard views are photographed the records are sufficiently comprehensive.

The electrokymograph is a device for securing an accurate graphic record of pulsation at any point on the cardiac border (Henny and Boone



Fig. 68—Normal kymogram (A.P. view)
(By courtesy of Dr. Jen et al.)

1947) A photosensitive pick up unit is placed between the patient and the screen so that the lead slit aperture lies across the border of the heart at the point where it is desired to record pulsation. The amount of light transmitted through the aperture varies with the movements of the cardiac border and is recorded graphically by means of a galvanometer operated by the photo electric cell. The interpretation of the graph is assisted by a simultaneous jugular phlebogram or other reference tracing.

ANGIOCARDIOGRAPHY

Individual cardiac chambers may be visualised by introducing 20 to 60 ml. of 70 per cent aqueous solution of diodrast or other contrast media (e.g. diodone) intravenously (Robb and Steinberg 1938) its course being recorded by means of serial skiagrams taken at intervals of one half to one second with the aid of a rapid cassette changer (Sussman, Steinberg and Grishman 1941) or serial fluorophotographs may be obtained by means of a special camera or cinematograph (Stewart, Breimer and Maier 1941). Contrast outlines of the right and left sides of the heart may thus be studied separately.

Patients may be seated or lying down. A long cannula of 6 to 9 inch length of wide bore plastic tubing is inserted into the antecubital vein and tied in position. A 50 ml. syringe is used and the diodone is injected through the cannula as rapidly as possible—usually within two or three seconds according to the volume. If a special camera, multiple cassette changer or cinematograph is not available, good pictures of the right side of the heart (fig. 2.09) may be obtained by exposing a film the instant the plunger is home. To obtain a satisfactory skiagram of diodone in the left side of the heart (fig. 2.10) without a special photographic unit is more difficult, but the technique is facilitated by measuring the circulation time beforehand. If 5 ml. of saccharine or decholine is used for this purpose the time obtained will prove too long to record the left sided angiocardio-gram for the large volume of diodone injected increases the speed of the circulation considerably. It has been found by experience that this momentary acceleration almost halves the circulation time. Due allowance should be made for this when calculating the best time to expose the film. Alternatively, however, the same cannula and the same volume of fluid used for angiocardio-graphy may be employed for measuring the circulation time, diluted saccharine giving a satisfactory end point. Under these circumstances the times should be identical.

The method has already been of service in determining the nature of the bump between the pulmonary artery and left ventricle in cases of mitral stenosis, showing it to represent the left auricle rather than the conus of the right ventricle (Robb and Steinberg 1939, Grishman, Sussman and Steinberg 1944) in establishing the diagnosis of certain congenital shunts, e.g. Fallot's tetralogy (Stewart, Breimer and Maier, 1941), patent ductus (Steinberg, Grishman and Sussman 1941), severe pulmonary stenosis



(a) Antero posterior view



(b) Second oblique position

Fig 2 09—Normal angiogram of the right side of the heart



(a) Antero posterior vie



(b) Second obl que position

Fig 2 10—Normal angiogram of the left side of the heart

with atrial septal defect Eisenmenger's complex, and tricuspid atresia in demonstrating coarctation of the aorta (Grishman Steinberg and Sussman 1941) and partial or complete superior vena cava obstruction (Taylor and Shulman 1942) and in distinguishing aneurysms of the aorta or pulmonary artery from other mediastinal masses (Thompson 1941)

CARDIAC MEASUREMENTS

Numerous measurements have been elaborated to serve as indices of enlargement of the heart or of one or more of its chambers but they do not compare with expert opinion based on the methods already outlined. The most reliable is the cardio thoracic ratio which is the transverse

diameter of the heart (fig 2 11) over the widest internal diameter of the thorax and which should not exceed 0.5. In normal adults the transverse diameter of the heart averages 12.2 cm in the male and 11 cm in the female the range being 8 to 14.5 cm (Roesler 1937)

The long diameter is measured from the junction of the superior vena cava and right auricle to the apex of the left ventricle and lies between 10 and 15.5 cm averaging 13 cm (Roesler 1937). It is especially increased in cases of left ventricular enlargement but it is also relatively increased in the long narrow heart of asthenic subjects.

The broad diameter is the sum of two perpendiculars drawn from the long diameter to the right cardio phrenic angle below and to the point of opposing movement on the left border of the heart above and measures 7 to 11 cm in normal adults with an average of 9 cm (Roesler 1937). It may be increased in cases of mitral stenosis and pulmonary heart disease when the transverse and long diameters are normal.

The location of the point of opposing movement is important for it tends to be raised or lowered according to whether enlargement is mainly left or right ventricular respectively. Similar significance is attached to the length of the chord which subtends the arc of the left ventricle measured from the point of opposing movement to the left cardio phrenic angle this line is normally 6 to 12.5 cm long and averages 9 cm (Roesler 1937).

The antero posterior diameter of the heart is measured from telerao grams taken in the lateral position and varies between 7 and 11 cm, with an average of 9 cm. It is a useful check on the significance of an increased transverse diameter for if this is due to cardiac enlargement the antero

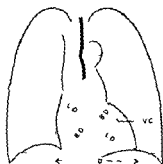


Fig 2 11—Diagram showing the common cardiac measurements

TD	Trans d m t
BD	Broad d m r
A	Width f rt
LD	Long d m t
LVCL	Let nter ul ho d

posterior diameter should be increased proportionately whereas if it is due to *depression of the sternum* the depth of the heart is decreased. The antero posterior diameter is especially increased in mitral stenosis.

The width of the aorta (2 to 3 cm) may be measured in the antero posterior or oblique positions whichever presents the clearest view of two sides of the vessel. In the antero posterior view the measurement should be made from the left side of the barium filled œsophagus to the left border of the aortic knuckle but it is only valid when the posterior part of the aortic arch passes directly backwards i.e. in a direction perpendicular to the frontal plane. In the oblique views barium in the œsophagus may also be helpful in the second oblique position for example the œsophagus may be deflected abruptly as it crosses the aorta so that the width of the vessel is seen clearly. In practice a normal aorta is most easily measured in the antero posterior view a syphilitic atheromatous or unfolded aorta in the second oblique view.

NORMAL VARIATIONS

Both the size and shape of the heart vary greatly in normal individuals thus in children and adolescents the pulmonary artery may be relatively prominent (fig. 2 12) in lean asthenic subjects the heart may be elongated and central in position (fig. 2 13) in short stocky individuals it is apt to lie transversely (fig. 2 14).

Displacement or rotation of the heart to left or right is often due to scoliosis the common finding being displacement of the heart to the left the spinal curvature being convex to the right. Rotation of the spine without conspicuous lateral curvature may cause considerable displacement or rotation of the heart. When cardiac displacement is due to partial collapse of the lung increased translucency of the over expanded normal lung on the same side is usually observed and is a valuable sign when the collapsed part cannot be seen.

Slight enlargement particularly of the left ventricle and of the transverse diameter is often seen in patients with slow heart rates whether due to sinus bradycardia sino auricular block or to heart block. The enlargement depends upon increased diastolic filling the slow rate being compensated by a large stroke volume (fig. 2 15). Slight enlargement of similar type may be encountered in athletes in some it may be explained by sinus bradycardia which is common in these subjects but in others it may be due to the extra demands which have been made on the heart.

In obese subjects the left cardio phrenic angle may be filled out by a triangular pad of fat (fig. 2 16) this must not be confused with left ventricular enlargement. In cases of depressed sternum the antero posterior skiagram may reveal general enlargement of the heart shadow but in the oblique views the depth of the heart is seen to be correspondingly reduced (fig. 2 17).

When such causes can be excluded and unsuspected enlargement of the

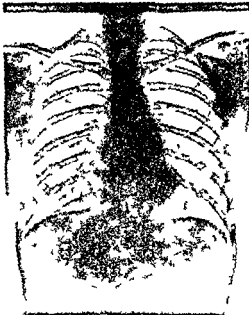


Fig. 212—Typical position of a relatively high heart in a normal subject



Fig. 213—Transversely placed heart of a heart block subject



Fig. 214—The elevated, relatively placed heart of a lean athletic subject



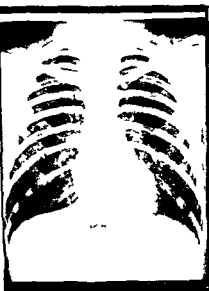
Fig. 215—Relatively large diastolic heart size in a subject with sinus bradycardia



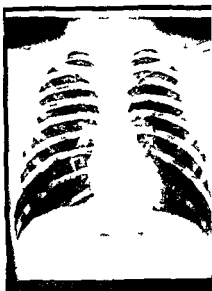
Fig 2 16—Teleradiogram of an obese subject showing a triangular opacity at the apex of the heart (pericardial fat)



Fig 2 17—Apparent enlargement of the heart in a case of depressed sternum



(a) The heart in diastole



(b) The heart in systole

Fig 2 18—Teleradiograms of the same patient taken with short exposure showing difference in size of heart shown in diastole and systole

cardiac silhouette is revealed by a skiagram it is wise to check the technique employed. Portable X rays or pictures taken with the patient lying or sitting may be misleading owing to distortion. Short exposures may catch the heart in systole and a skiagram so obtained may be appreciably smaller than one photographed in diastole (fig 2 18)

The heart may be smaller than normal in many wasting diseases when atrophy takes place but this is of little practical importance

RADIOGRAPHIC ABNORMALITIES

The illustrations referable to this section may be found in other chapters but for the sake of convenience are also reproduced here

ABNORMALITIES OF THE AORTA

Saccular aneurysm (fig 2 19 a and b) is pathognomonic of syphilis. It may be distinguished from other space filling lesions by its intimate connexion with the aorta in all views by calcification of its walls and by its



Fig 2 19—Saccular aneurysm of the aorta
(a) Anteroposterior view (b) Second oblique position
(By courtesy of S. John Parkinson)

pulsation a thrombosed sac however may not pulsate. Angiocardiography is helpful in doubtful cases. Fusiform aneurysm (fig 2 20) is usually associated with aortic incompetence and is also practically diagnostic of syphilis. It should be distinguished from undue prominence of the aorta such as that commonly found in other forms of aortic incompetence (fig

221) occasionally it is due to dissection associated with cystic medial necrosis of the aorta. Syphilitic aortitis without aneurysm or fusiform dilation cannot be diagnosed radiologically with assurance although irregularities of calibre if clearly demonstrable are suggestive.

Unfolding of the aorta may occur in aortic valve disease in hypertensive heart disease and in atherosclerosis. The ascending limb is conspicuous the knuckle is unduly prominent and the descending limb appears to the



FIG. 220 (a)—Fusiform aneurysm of the aorta (anteroposterior view)



FIG. 220 (b)—Fusiform aneurysm of the aorta (2nd oblique position)

patient's left in the anteroposterior view (fig. 222). In the second oblique position the arch is wider than normal and its posterior part may pull the oesophagus backwards (fig. 223). Vigorous pulsation proclaims aortic incompetence rather than hypertension or atherosclerosis.

Tortuosity of the aorta is characteristic of atherosclerosis. It is best seen in the second oblique view but may be so marked that the descending limb appears to the right of the heart shadow in the anteroposterior view (fig. 224). Calcification of the aorta (fig. 225) is also characteristic of atherosclerosis but may occur in the wall of a syphilitic aneurysm.

Coarctation of the aorta (fig. 226) may be recognised by a relatively small absent or elongated aortic knuckle by notching of the ribs and by the angiocardigraphic demonstration of the constriction itself. In addition

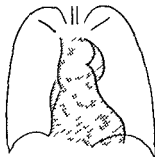


Fig. 224 — Orthodiagram illustrating tortuosity of the aorta



Fig 2 21—Prominence of the aorta due to rheumatic aortic incompetence



Fig 2 22—Unfolding of the aorta in hypertensive heart disease



Fig 2 23—Unfolding of the aortic arch illustrated by barium in the esophagus



Fig 2 25—Calcification of the aortic arch

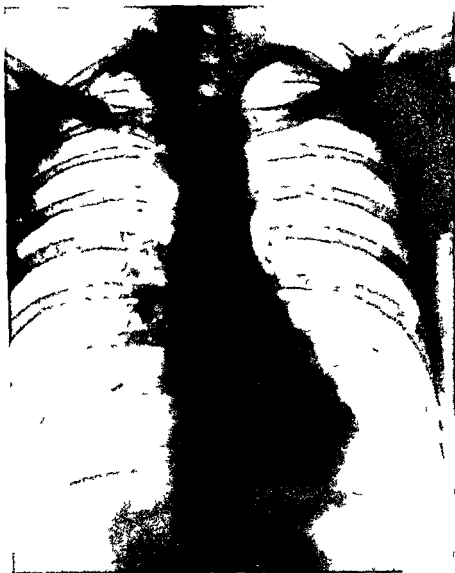


Fig 2 6—Coarctation of the aorta showing hypoplasia of the aorta and notching of the inferior margins of the ribs

the left ventricle is usually enlarged. Absence of the aortic knuckle is also associated with congenital right sided aorta but the latter may be distinguished by the behaviour of the barium filled œsophagus which is then deflected to the patient's left instead of to his right as it crosses the aorta (fig 2 27)

Hypoplasia of the aorta is rare as a solitary congenital abnormality but is common in association with certain other congenital or acquired lesions especially atrial septal defect and mitral stenosis. The aortic knuckle is small and its pulsation diminished (fig 2 28)



(a) Antero posterior view



(b) First oblique position

Fig 2 27 —Right sided aortic arch illustrated by means of barium in the œsophagus

(H. ou t j f S J h Park)

ABNORMALITIES OF THE LEFT VENTRICLE

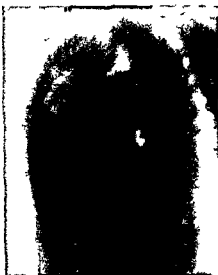
Left ventricular enlargement is encountered chiefly in hypertensive heart disease and aortic valve disease but it is also seen in patent ductus arteriosus and in organic mitral incompetence and may occur in various conditions as part of general enlargement. It is easily recognised by the density and bulk of the left ventricular shadow in the antero posterior and second oblique positions by increase in the transverse and long diameters of the heart and of the left ventricular chord and by elevation of the point of opposing movement (figs 2 29a and 2 29b). In hypertension and aortic valve disease the shadows of the unfolded aorta and of the heart itself may be compared either to two ovals set at right angles or to the shape of a boot



Fig 228—Hypoplasia of the aorta in a case of mitral stenosis



(a) Antero posterior view



(b) Second oblique position

Fig 229—Enlargement of the left ventricle due to syphilitic aortic incompetence

When there is left ventricular failure the hilar shadows are exaggerated representing engorged pulmonary vessels (fig 2 29a) and there may be fan shaped mottling spreading outward towards the periphery if there is pulmonary œdema (fig 2 30) Hydrothorax may be present and if unilateral is usually left sided (Bedford and Lovibond 1941)

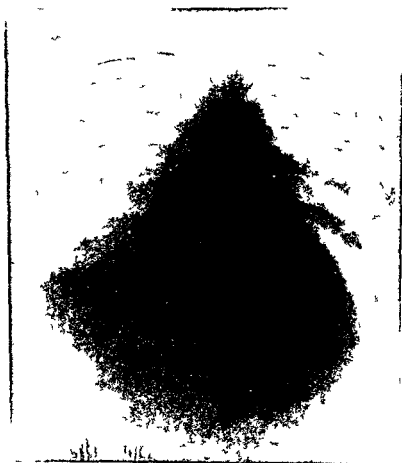


Fig 2 30—Acute pulmonary œdema due to left ventricular failure in a case of malignant hypertension

Left ventricular aneurysm may present as a bulge on the left border of the heart, usually towards the apex (fig 2 31) and may exhibit paradoxical pulsation. Myocardial infarction may be located with precision in some cases by the fluoroscopic demonstration of an area with absent or paradoxical pulsation. This can be well shown by kymography (fig 2 32)

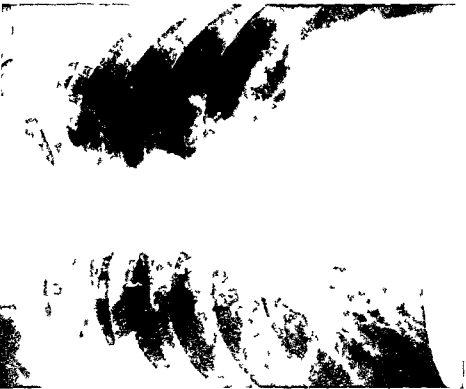


Fig 2 31—Left ventricular aneurysm

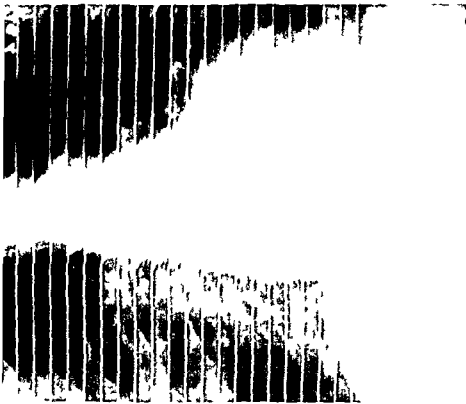


Fig 2 32—Kymogram showing absence of pulsation over an anterior lateral aneurysm of the left ventricle
(H 1007 H 1008)



Fig. 23.—Dilatation of the left auricle forming a bump between the pulmonary arc and left ventricle in a case of organic mitral incompetence

DILATATION OF THE LEFT AURICLE

Conspicuous dilatation of the left auricle invariably means organic mitral valve disease but the chamber may be unduly full in cases of left ventricular failure. In the antero posterior view it may appear as a bump on the left border of the heart between the pulmonary artery and left ventricle (fig 2 33) where it has been mistaken for the conus of the right ventricle. The best proof that this bump represents a dilated left auricle is afforded by

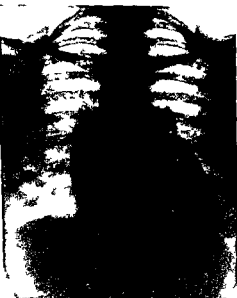


Fig 2 34—Dilatation of the left auricle seen on both borders of the heart in the antero posterior view in a case of mitral stenosis



Fig 2 37—Dilatation of the left auricle seen in the 2nd oblique position with barium in the œsophagus. Case of mitral stenosis

angiocardiology (Grishman Sussman and Steinberg 1944) by passing a catheter into the area via a patent foramen ovale or by observing its pulsation in cases of heart block or mitral incompetence. On the right border of the heart an enlarged left auricle may throw a convex shadow above but overlapping that of the right auricle (fig 2 34). The barium filled œsophagus is usually deflected to the patient's right in the antero posterior view.

In the right anterior oblique position the œsophagus is displaced backwards in an abrupt manner immediately below the left bronchus and pulmonary artery (fig 2 35) the antero posterior diameter of the heart being increased and the retrocardiac space decreased correspondingly. Backward displacement of the œsophagus from an enlarged left ventricle is rarely so abrupt or so high but on occasions it may be indistinguishable (fig 2 36). In the left anterior oblique position an enlarged left auricle

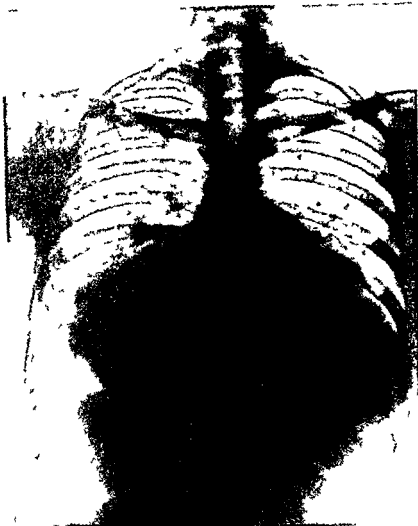


Fig 278 (a)—Aneurysmal dilatation of the left auricle in a case of mitral stenosis. Antero posterior view



Fig. 238 (b)—Aneurysmal dilatation of the left auricle in same case of mitral stenosis. First oblique position.

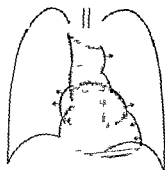


FIG. 2 39—Orthodiagram illustrating expansion of the left auricle during ventricular systole in a case of organic mitral stenosis

causes the œsophagus to be deflected backwards above the shadow of the left ventricle (fig 2 37)

Aneurysmal dilatation of the left auricle (fig 2 38 a and b) may be caused by rheumatic mitral incompetence or stenosis but it is probable that a high degree of auricular muscle damage is an important contributory factor

Systolic expansile pulsation of the left auricle is pathognomonic of mitral incompetence usually organic. It is especially convincing when seen on both borders of the heart in the antero posterior view (fig 2 39). In the first oblique position minor degrees of backward pulsation of the left auricle are often seen and must be regarded as normal but the quality and amplitude of the movement in organic mitral incompetence are most impressive and are easily recognised with experience. Electrokymographic tracings should prove helpful in doubtful cases.

ABNORMALITIES OF THE PULMONARY ARTERY

Dilatation of the pulmonary artery occurs chiefly in three congenital and in three acquired lesions namely patent ductus arteriosus atrial septal defect and pulmonary stenosis with closed septa on the one hand and mitral stenosis pulmonary heart disease and beri beri on the other. Rare causes include the Eisenmenger complex and capricious cases of patent interventricular septum without a dextraposed aorta sometimes no explanation is found.

In patent ductus arteriosus (fig 2 40) dilatation and exaggerated pulsation of the pulmonary artery and its proximal branches are usually associated with a hyperkinetic aorta enlargement of the left ventricle and fullness of the left auricle. Angiocardiography may reveal a shadow which resembles a small aneurysm of the inferior margin of the aortic arch where it is joined by the ductus (Steinberg *et al* 1943). Atrial septal defect (fig 2 41) is characterised by gross dilatation and conspicuous pulsation of the pulmonary artery and its two main branches by a heavily marked peripheral pulmonary vascular tree by considerable enlargement of the right ventricle and right auricle especially if there is associated mitral stenosis (Lutembacher's syndrome) and by hypoplasia of the aorta (Bedford Papp and Parkinson 1941). In pulmonary stenosis with closed septa dilatation and visible pulsation of the pulmonary artery are more or less confined to the main vessel (fig 2 42) and are slight or moderate in degree the right ventricle is enlarged but not the auricle (unless there is failure). In Eisenmenger's complex considerable dilatation of the pulmonary artery

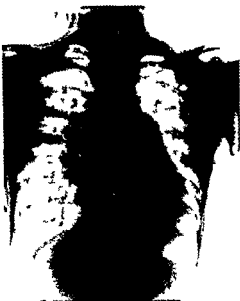


Fig. 2 40—Dilatation of the pulmonary artery and its branches associated with left ventricular enlargement due to patent ductus



Fig. 2 42—Dilatation of the pulmonary artery in a case of pure pulmonary stenosis



Fig. 2 43—Dilatation of the pulmonary artery in a case of Eisenmenger's complex
(By courtesy of Dr. Mac Campbell)



Fig. 2 44—Dilatation of the pulmonary artery associated with pulmonary congestion in a case of mitral stenosis



Fig. 2.41.—Dilatation of the pulmonary artery and its branches associated with hypoplasia of the aorta and right ventricular enlargement in a case of atrial septal defect



(a) Antero-posterior view



(b) First oblique position

Fig 2 45—Dilatation of the pulmonary artery and its main branches in a case of chronic pulmonary heart disease due to emphysema



Fig 2 45 (c)—Second oblique position



Fig 2 46—Dilatation of the pulmonary artery due to primary or idiopathic pulmonary hypertension

is the rule (fig 2 43) as the shunt is from right to left the aorta is not hypoplastic and the peripheral pulmonary vascular shadows are inconspicuous. The appearances in ventricular septal defect resemble those of patent ductus but the pulmonary artery is usually less dilated.

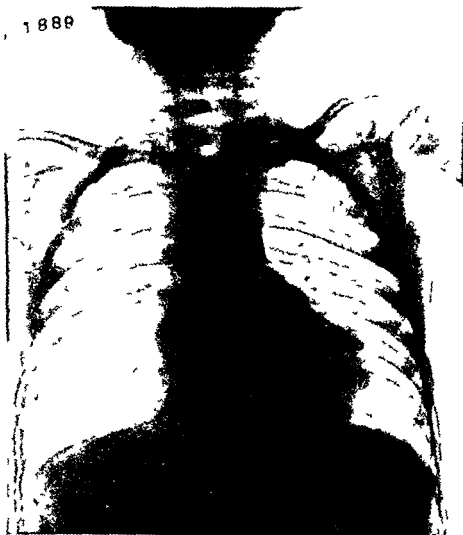


Fig 47—The *Cœur en sabot* due to Fallot's tetralogy

Of the acquired lesions mitral stenosis may usually be recognised by the presence of aortic hypoplasia and enlargement of the left auricle but these are not always evident (fig 2 44) moreover if there is aortic incompetence as well the aorta may be prominent and the left ventricle enlarged. If there is a bump between the pulmonary artery and left ventricle in the antero-posterior view mitral stenosis (or incompetence) is certain whether the shadow is believed to represent the left auricle or the conus of the right

ventricle for it does not occur in any other condition. In pulmonary heart disease the pulmonary artery and its main branches are moderately prominent (fig 2.45a). Owing to associated emphysema the rounded shadow of the bifurcation of the pulmonary artery appears especially dense by contrast in the first oblique position (fig 2.45b) and the pulmonary arc is unusually well defined even to dwarfing the aortic arch in the second oblique position (fig 2.45c). The aorta is more prominent in the antero-posterior view than in the other congenital and acquired lesions mentioned above owing to the high cardiac output. The right ventricle is enlarged and the right auricle is abnormally full. Pulmonary heart disease without emphysema, so called primary pulmonary vascular sclerosis (fig 2.46) or idiopathic pulmonary hypertension is more easily confused with congenital lesions especially with simple pulmonary stenosis. Pure beri beri is rare in the United Kingdom but is not uncommon in Japan, China, Java and other eastern countries. The radiological appearances are similar to those of anoxic pulmonary heart disease but there is no emphysema.

Hypoplasia of the pulmonary artery is characteristic of Fallot's tetralogy (fig 2.47). There may be a distinct gap between the aortic knuckle and the curve of the left ventricle and the vascular shadows at the hilum are reduced on both sides.

ENLARGEMENT OF THE RIGHT VENTRICLE

Right ventricular enlargement is more difficult to recognise than left. In the antero-posterior view there is usually some increase in the transverse and broad diameters, the right auricle being pushed a little to the right and inter-ventricular septum to the left. In some cases the septum is so far to the left that it forms the left border of the heart, only the tip of the left ventricle being visible from the front; the effect produced is that of increased angularity of the cardiac apex and a more acute left cardiophrenic angle, the general shape resembling the Dutch peasant's wooden shoe with turned up toe. This is the *cœur en sabot* and is especially characteristic of Fallot's tetralogy (fig 2.47).

In the left anterior oblique



Fig 2.48—Right ventricular enlargement in a case of mitral stenosis (2nd oblique position)



Fig 2 49—Cross enlargement of the right auricle due to tricuspid incompetence

position right ventricular dominance is recognised by the increased curvature of the anterior border of the heart shadow. Instead of the lob-sided appearance resulting from normal left ventricular bias the heart shadow is more globular the anterior and posterior ventricular curves being more equal (fig 2 48). If the right auricle is enlarged however as may be determined from the antero posterior view interpretation is more difficult for its shadow is superimposed on that of the right ventricle in the second oblique position and it may be entirely responsible for the increased curvature of the anterior border.

The right ventricle is disproportionately enlarged in Fallot's tetralogy and in all conditions associated with dilatation of the pulmonary artery except patent ductus and patent interventricular septum.

ENLARGEMENT OF THE RIGHT AURICLE

Dilatation of the right auricle usually associated with fullness of the superior vena cava is seen in most cases of congestive heart failure and of atrial septal defect but it reaches gross proportions in tricuspid stenosis or incompetence and in Lutembacher's syndrome. It may be distinguished from pericardial effusion by the blunt right cardio-phrenic angle (cf figs 2 49 and 2 50).

When the shadow of a dilated left auricle appears on the right border of the heart it should be recognised by its higher position and more abrupt curvature (fig 2 34). A zone of increased density is seen where it overlaps the shadow of the right auricle. A grossly tortuous aorta occasionally gives rise to confusion for its descending limb may appear to the right (fig 2 24). Fluoroscopy in the oblique positions should clarify the issue.

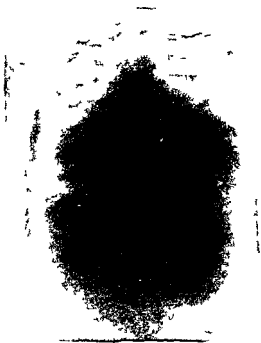


Fig 50—Pericardial effusion showing acute right cardio-phrenic angle



Fig 251—General enlargement of the heart due to isolated myocardia

GENERAL ENLARGEMENT OF THE HEART

General enlargement of the heart shadow involving all diameters is seen in rheumatic diphtheritic and Fiedler's carditis (fig 2 51) in anæmia thyrotoxicosis arteriovenous aneurysm and extensive active Paget's disease of bone in myxœdema and sometimes in von Gierke's disease in rheumatic heart disease with multiple valve lesions and in fibrosis of the myocardium of uncertain etiology. An example of general enlargement due to thyrotoxic heart failure is shown in fig 2 52

PERICARDIAL EFFUSION

When effusion is considerable the natural contours of the heart in the antero posterior view are replaced by single bold curves and pulsation is absent or greatly reduced. The right cardio phrenic angle is unusually acute (fig 2 50). In the first oblique position the posterior border of the heart shadow may bulge beyond the barium filled œsophagus. When effusion is slight or moderate however radiological diagnosis may be difficult for pulsation may be clearly visible and may indeed be greater than that often seen in some of the conditions most likely to cause confusion. Moreover the natural contours of the heart on one or other border may not be entirely lost and may closely imitate the indefinite demarcations of the chambers which may be seen in acute carditis and myxœdema for example. Under these circumstances more reliance should be placed on change of shape with alteration of posture on bulging of the posterior inferior angle of the heart shadow in the first oblique position (instead of the normal concavity of the inferior vena cava) and on the degree and rapidity of changes in size from week to week.

CONSTRUCTIVE PERICARDITIS

The most important radiological evidence of constrictive pericarditis is loss of pulsation without cardiac enlargement but calcification of the pericardium is common and helpful and is usually best seen in the left anterior oblique position (fig 2 53). Slight to moderate enlargement of the heart shadow may occur if the pericardium is sufficiently thick (1 to 2 cm) but the triangular appearance given by the obliquely set straight right and left borders should suggest the correct diagnosis (fig 2 54).

CALCIFIED VALVES

Calcified valves are best seen fluoroscopically they may be recorded by means of tomography. The patient should be turned 15 degrees to the left and an imaginary line drawn from the point of opposing movement on the left border of the heart downwards and to the patient's right at an angle of 45 degrees with the horizontal (fig 2 55). The aortic valve is situated just above this line in the centre of the heart shadow the mitral just below



Fig 252—General enlargement of the heart due to thyrotoxic heart failure with auricular fibrillation



Fig 253—Case of chronic constrictive pericarditis showing extensive calcification of the pericardium (2nd oblique position)



Fig 254—Chronic constrictive pericarditis showing triangular shaped heart in the antero-posterior view

it and a little to the patient's left. Calcification may be recognised by linear or anti clockwise elliptical movement of dense crescentic opacities in the direction of the anatomical axis of the heart synchronous with the heart

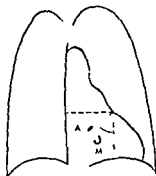


Fig. 255—Orthodiagram showing the position of calcified valves. The patient has been turned fifteen degrees to his left.

A A M I
M M I I

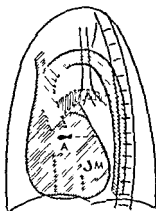


Fig. 256—Orthodiagram showing the position of calcified valves in the 2nd oblique position.

A A M I
M M I I

beat. The technique requires proper accommodation and maximum constriction of the diaphragm so that only a square inch or so of the screen is visible. Calcified aortic valves are sometimes better seen in the second oblique position where they lie at the intersection of a vertical line through the centre of the heart shadow and a horizontal line through the top of the left ventricular arc (fig. 256). This view may be helpful in valve differentiation for the mitral valve lies in the posterior third of the heart shadow (Sosman, 1939) and at a lower level.

REFERENCES

- Bedford D. E. and Lovibond J. L. (1941) Hydrothorax in heart failure. *Brit Heart J* 3 93. — Papp C. and Parkinson J. (1941) Atrial septal defect. *Ibid* 3 37.
- Grishman A., Steinberg M. F. and Sussman M. L. (1941) Tetralogy of Fallot: contrast visualisation of the heart and great vessels. *Radiology* 37 178. — Sussman M. L. and Steinberg M. F. (1944) Angiocardiographic analysis of the cardiac configuration in rheumatic mitral disease. *Amer J Roentgenol* 51 33.
- Henny G. C. and Boone B. R. (1947) Improved electrokymograph for recording heart motion: improved type. *Ibid* 57 409.
- Robb G. P. and Steinberg I. (1938) A practical method of visualisation of the chambers of the heart, the pulmonary circulation and the great vessels in man. *J clin Invest* 17 507. — (1939) Visualisation of the chambers of the heart. *Amer J Roentgenol* 42 14.

Roesler H (1928) Beiträge zur Lehre von den angeborenen Herzfehlern
IV Untersuchungen an zwei Fällen von Isthmus Stenose der Aorta *Weener
Arch inn Med* 15 521 — (1937) Clinical roentgenology of the cardiovascular
system London

Sosman M C (1939) Roentgenological aspects of acquired valvular heart
disease *Amer J Roentgenol* 42 47

Steinberg M F Grishman A and Sussman M L (1943) Angiocardiography in congenital heart disease II Intracardiac shunts *Ibid* 49 766

III Patent ductus arteriosus *Ibid* 50 306

Stewart W H Breimer C W and Maier H C (1941) Cine-roentgenographic diagnosis of congenital and acquired heart disease *Ibid* 46 636

Stumpf P (1931) Archiv und Atlas der normalen und pathologischen Anatomie in typischen Röntgenbildern Das röntgenographische Bewegungsbild und seine Anwendung (Flächenkymographie und Kymoskopie) Fortschr a d Geb d Rongenstrahlen [Ergänzungsband 41 Georg Thieme Leipzig]

Sussman M L Steinberg M F and Grishman A (1941) Multiple exposure technique in contrast visualisation of the cardiac chambers and great vessels *Amer J Roentgenol* 46 745

Taylor H K and Shulman I (1942) Cardio angiography *Radiology* 39 523

Thompson S A (1941) Differential diagnosis by means of intravenous contrast medium of two cases simulating aneurysm of the pulmonary artery *Amer J Roentgenol* 46 646

ELECTROCARDIOGRAPHY

ELECTROCARDIOGRAPHY was discovered in relation to the frog's heart by Kolliker and Müller (1856) and was proved applicable to the study of the heart in man by Waller (1887) who used a capillary electrometer and an antero posterior chest lead. It was elaborated by Einthoven (1903) inventor of the string galvanometer and author of the famous triangle which bears his name and used extensively by Lewis (1925) in his well known researches on abnormalities of rhythm. In recent years many attempts have been made to place electrocardiography upon a more scientific and less empirical basis and considerable success has been achieved in this respect especially by Wilson and his colleagues (1930 *et seq.*) It is not easy (or necessary) for the ordinary physician unless he also be a physicist and mathematician to grasp the electrical details involved but the following simplified account will be readily understood.

Certain molecules in the resting cardiac muscle cell dissociate into positive and negative ions. The positively charged ions (cations) are distributed on the outer surface the negatively charged ions (anions) within (Curtis and Cole 1941). Such a cell is in a state of electrical balance and is said to be polarised (fig. 3 01a). When the cell is excited its polarity is reversed the negative charges coming to the surface the positive charges passing within and the cell is said to be depolarised (fig. 3 01b). It should be clear that when a number of cells are clustered together all in the resting polarised state or all in the excited depolarised state there can be no potential differences anywhere on their collective surface. If a group of cells were in the process of being excited however those already depolarised would possess negative surface charges whereas those still polarised would have positive surface charges and the collective surfaces of the two sets would yield a potential difference (fig. 3 01c). This constitutes a doublet (Craib 1930) dipole (Ashman 1948) or double layer (Bayley 1943). Thus when an excitatory wave flows through cardiac muscle its head is electrically positive and its tail negative (fig. 3 01d). If electrodes are placed at A and B and connected to a galvanometer an electrical current flows from B to A through the galvanometer and from A to B through the tissue. The excitatory process or accession wave as it is called causes a very rapid or almost instantaneous reversal of cellular polarity so that the duration of the galvanometric deflection is brief and practically indicates the speed of the wave if the muscle thickness is known or the muscle thickness if the speed of the wave is known. When the impulse reaches B (fig. 3 01e) the whole muscle block AB has a negative collective

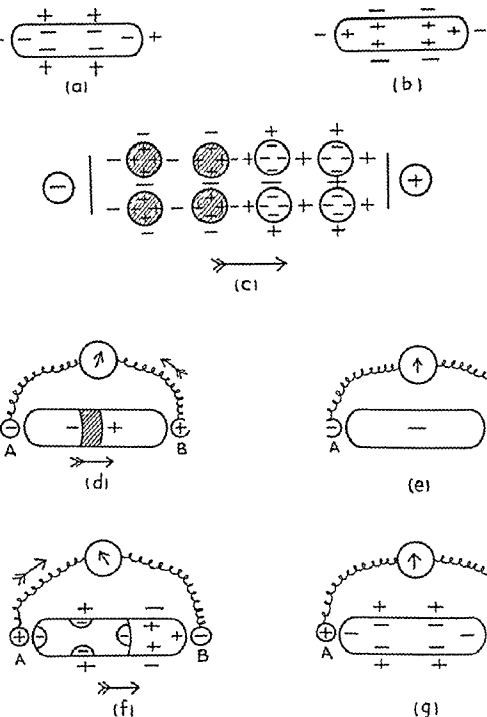


Fig 301—Accession and regression waves (stages of excitation and recovery) in cardiac muscles (see text)

- (a) Resting polarised cell
 (b) Excited depolarised cell
 (c) Excitation proceeding through a group of cells
 (d) Spread of the accession wave through a block of cardiac muscle
 (e) Muscle block fully excited (depolarised)
 (f) Spread of the regression wave or recovery process
 (g) Muscle block fully recovered (re polarised)

surface if recovery has not yet commenced at A and there is no potential difference between A and B. Within a short time however recovery begins at A (fig. 3 01f) and the cells become repolarised their collective surfaces becoming positively charged again. While the recovery process or regression wave as it is called is spreading from A towards B a current again flows through the galvanometer but in the opposite direction. The regression wave travels at the same speed as the accession wave but causes a slower change of polarity so that the galvanometric deflection is not so brief. If the movements of the galvanometer are graphically recorded the passage of an excitatory impulse from A to B results therefore in a diphasic curve such as that shown in figure 3 02 the first deflection being quick or sharp the second slow or blunt. Moreover if the neuro muscular tissue is uniform in all relevant respects the area occupied by the first deflection which may be measured by means of a planimeter with suitable magnification is exactly equal though of opposite sign to the area occupied by the second deflection. In modern electrocardiographic parlance the first deflection is represented by the P wave when it reflects auricular excitation and

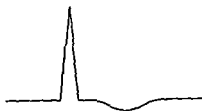


Fig. 3 02—The diphasic curve produced by the processes of excitation and recovery in heart muscle

by the QRS complex when it reflects ventricular excitation while the second is represented by the T_a and T_v waves respectively. The QRS complex is written as the accession wave flows through the heart muscle from endocardial to epicardial surfaces not as the excitatory impulse passes down the bundle of His bundle branches and Purkinje network. As the heart is not a uniform muscle block but a bi ventricular organ composed of numerous intertwining S shaped muscle bundles (Robb and Robb 1938) the initial ventricular deflection (QRS) is not monophasic as in figure 3 02 but complex and usually biphasic or triphasic nor is the second ventricular deflection (T) of equal area and opposite sign. On account of this complexity it is impossible in the light of present knowledge to determine by scientific theory precisely what an electrocardiogram should look like it is only possible to find out by the practical method. For this reason electrocardiography has largely remained an empirical study.

Einthoven's string galvanometer consists of an exceedingly fine fibre such as silver coated glass suspended between the poles of an electro magnet when a current passes through the fibre the latter is deflected towards one or other pole according to the direction of the current. By suitable magnification and illumination the movements of the shadow of this string may be recorded on a moving photographic film. Valve amplifying oscillographs of various form operated by potential differences may be used instead of Einthoven's instrument. Time marking is so arranged that fine vertical lines appear on the film at intervals of

0.04-0.05 second preferably with thicker lines every 0.20 second Horizontal lines for measuring voltage are spaced at intervals of 1 mm

Practical points to bear in mind include satisfactory insulation of the machine and lead wires to prevent 50 cycle A C interference proper standardisation of the galvanometer so that a deflection of 2 cm represents a potential difference of 1 mv and the elimination of skin resistance by means of electrode jelly The paste described by Jenks and Graybiel (1935) has proved effective it consists of sodium chloride 2950 G (6.5 lb) powdered pumice 3600 G (8 lb) gum tragacanth 226 G (8 oz) potassium bitartrate 114 G (4 oz) glycerol 710 ml (4.4 oz) phenol 28.5 G (1 oz) and water to 7.5 litres (2 gallons) The electrolytes are dissolved in one gallon of water while the gum and glycerol are heated for six hours in the other the two are then mixed stirred and reheated for one hour Phenol and pumice (and more water if necessary) are then added and mixed until the preparation has the consistency of cream Fresh soft green soap (B.P.) is very little inferior especially after rubbing the skin with some abrasive (Bell Knox and Small 1939) A number of satisfactory pastes or gels are marketed

CHLEST LEADS

Analysis of electrocardiograms has become simplified since the introduction of Wilson's neutral electrode (Wilson 1934) Previously all electrocardiograms were bipolar and registered the potential differences between two electrodes placed at different sites on the surface of the body each gathering different potential values According to Linthoven's theory however the algebraic sum of the potentials at the left arm right arm and left leg always equal zero these points representing the apices of an equilateral triangle in the frontal plane of the body the heart lying at its centre and the limbs being regarded as extensions of the lead wires * Thus it is only necessary to link up these three points to a common terminal (preferably through a resistance of 5,000 ohms in order to neutralise differences in skin resistance) to provide an electrode that remains at zero potential throughout the cardiac cycle If this neutral or indifferent electrode is linked to one arm of the galvanometer the instrument will record the potential variations of an exploring electrode linked to the other arm This is the basis of all V leads V standing for potential value or voltage at any particular point It has been agreed that positivity of this exploring electrode should be represented by an upright electrocardiographic deflection

It is now necessary to consider the variations in potential that may be recorded if the exploring electrode is placed over the surface of the left ventricle in man (Wilson *et al.* 1944) As the accession wave spreads from endocardial to epicardial surfaces the left ventricular cavity (in contact with the tail of the wave) becomes electrically negative and the surface of the heart (in contact with the head of the wave) becomes electrically positive The galvanometer therefore records an upright or positive deflection R (fig. 3.03b) When the accession wave reaches the surface the exploring

* The mathematical proof of this equation is given by Wilson *et al.* (1946) Goldberger (1947) and by others

electrode undergoes an abrupt reversal of polarity and the galvanometer registers a sharp downward deflection (the intrinsic deflection). As both the cavity and surface of the left ventricle are then at the same negative potential the electrical field is abolished and the galvanometer comes to rest (fig 3 03c). A complication arises however because the accession wave starts at some point (such as the left side of the interventricular septum) remote from the muscle underlying the electrode. The left ventricular cavity thus becomes negative before the muscle under the electrode

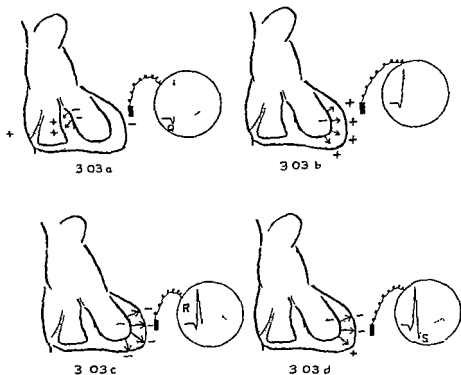


Fig 3 03—Formation of the Q R and S waves (see text)

begins to be activated and this negative potential is passively transmitted to the surface to be recorded as an initial downward deflection Q (fig 3 03a). As leads taken from the right ventricular cavity show an initial positive deflection in practically all instances it is now generally believed that the excitation wave starts on the left side of the septum. Again if the accession wave is still spreading through muscle remote from the exploring electrode when the galvanometer has registered the local intrinsic deflection the electrical field is maintained and continued negativity of the cavity is passively transmitted to the surface under the electrode to be recorded as a final downward deflection S (fig 3 03d).

When the exploring electrode is placed over the right ventricle similar principles hold good but the right ventricle is much thinner than the left

and therefore the local potential differences are smaller and are normally overpowered by left ventricular events. An initial R wave is almost invariable and probably represents the positive potential produced in the right ventricular cavity as the accession wave spreads through the septum from left to right. Further development of R as excitation passes through the anterior wall of the right ventricle is more or less prevented by the stronger negative potential induced by the tail of the accession wave that is spreading through the left ventricle; this is represented by a large S wave. Q is never seen over a normal right ventricle. The second ventricular deflection T is upright over the left ventricle but may be inverted over the right (in leads V₁ and V₂).

In clinical electrocardiography multiple chest leads are designated leads V₁-7. The figures indicate the position of the proximal electrode with reference to the chest wall and represent respectively the right and left

borders of the sternum, the left para-sternal and mid-clavicular lines and the anterior mid and posterior axillary lines at the level of a line passing from the fourth intercostal space at points 1 and 2 to the fifth intercostal space at point 4 and thence horizontally (fig 3 04) 1 or routine purpose leads V₁, V₃ and V₅ or V₂, V₄ and V₆ are usually sufficient but in particular instances other combinations or all seven leads are preferable. A typical record obtained with this technique is illustrated in fig 3 05. Over the left ventricle (V₅ and V₆) there is a small Q wave, a large R wave, no S wave

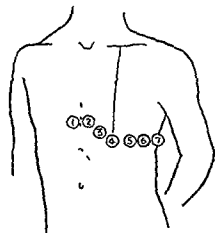


Fig. 3 04—Multiple chest leads V₁-V₆.
Position of the exploring electrode

an isopotential R-T junction and an upright T wave. In the transition zone (V₃-V₄) Q has disappeared, a conspicuous S wave has developed and T is sharply upright. Over the right ventricle (V₁) there is again no Q wave, R is small, S large and T is flattened. In normal subjects the P wave is upright or occasionally diphasic (3 per cent) in V₃ but often diphasic (20 per cent) or inverted (15 per cent) in V₁. Q is usually present in V₆, occurs in V₅ in 45 per cent of cases, but is rarely seen farther to the right. S is usually absent in V₆-7, is absent in V₅ in 17 per cent of cases but is invariably found in V₃ and V₁. T is always upright in V₄-V₆, may be occasionally diphasic in V₃ and is inverted in V₁ in 62 per cent of cases.

If there is clockwise rotation about the longitudinal axis (viewed from below) the anterior surface of the septum is shifted to the patient's left; this means that S is dominant in V₄, the transition zone being shifted to V₄ or V₅. In such cases Q may not appear until V₆ or V₇. Similar graphs

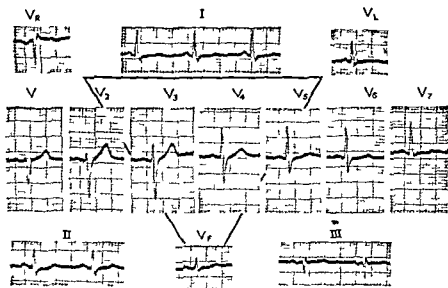
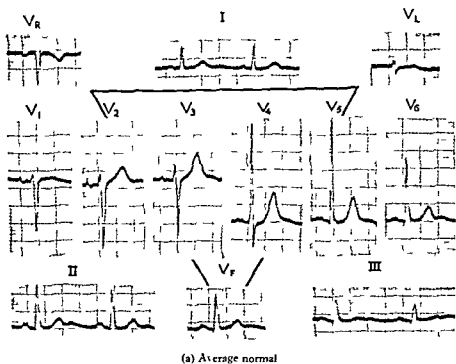


Fig 3.05 - Normal chest lead electrocardiogram (V_1 - V_6)

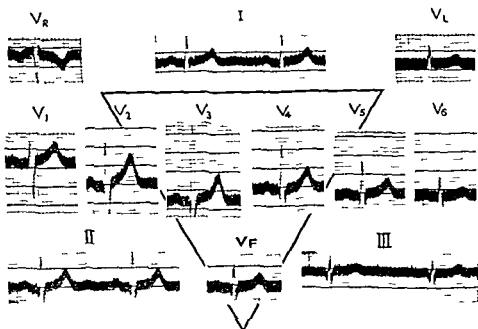


Fig 3 05 (c)—Anti clockwise rotation

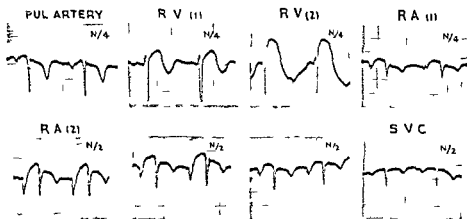


Fig 3 06—Right ventricular and pulmonary artery cavity leads

are obtained when the heart is horizontal in position the septum then being displaced to the patient's left (fig. 3 05b)

Anticlockwise rotation about the longitudinal axis brings the anterior surface of the septum to the patient's right. The QR pattern may then be seen from V₆ to V₃ and the transition zone is shifted to V₂ (fig. 3 05c)

In addition to leads V₁-V₇ other positions of the exploring electrode have been used with advantage under exceptional circumstances. An œsophageal lead may also be helpful in doubtful cases of posterior myocardial infarction and an intracardiac lead may provide interesting information but these are rarely necessary for clinical purposes.

The œsophageal lead takes its potential from the surface of the left auricle when high and from the posterior surface of the left ventricle when low. Left auricular potentials are transmitted from the cavity of the left ventricle and show monophasic Q waves and inverted T waves, the cavity of the left ventricle being negative throughout the inscription of the initial and second ventricular deflections. The posterior surface of the left ventricle gives rise to a QR complex similar to that obtained anteriorly or laterally. Œsophageal patterns therefore show monophasic Q waves or QR deflections, Q dominating when the electrode is relatively high up, R when the electrode is relatively low down. T is usually negative when the electrode is high, positive when low.

Intracardiac leads from the cavity of the right ventricle show a small initial R wave followed by a deep S wave as already described. If the catheter is passed through a patent foramen ovale into the left ventricle a monophasic Q wave is obtained. When the catheter is passed into the pulmonary artery the small R wave seen within the cavity of the right ventricle disappears in favour of a monophasic Q wave (fig. 3 06) this is because the pulmonary artery takes its potentials from the surface of the left auricle.

There are thus only a limited number of basic QRS patterns upon which all ventricular deflections encountered in clinical electrocardiography depend (fig. 3 07): the QR complex of a left ventricular surface lead (T normally upright); the RS complex of a right ventricular surface lead (T usually upright); the monophasic Q wave of a left ventricular cavity lead (T normally inverted); the RS complex of a right ventricular cavity lead (T normally inverted); and the balanced QR pattern of a combined left ventricular cavity and surface lead from the back of the heart (Goldberger 1947).

The direction of the second ventricular deflection T is opposite to theoretical prediction in all the basic patterns and suggests that the recovery wave starts at the surface of the ventricles and is directed towards the cavities.

Instead of V lead many workers including Wolferth and Wood (1932-33) who re-introduced chest leads to clinical electrocardiography have coupled the exploring electrode with a relatively indifferent electrode

placed on the right arm (CR) or on the left leg (CF) Agreement will never be reached as to which of these is the more informative and it is expected that they will both be abandoned in favour of V leads They will be considered in greater detail in subsequent paragraphs

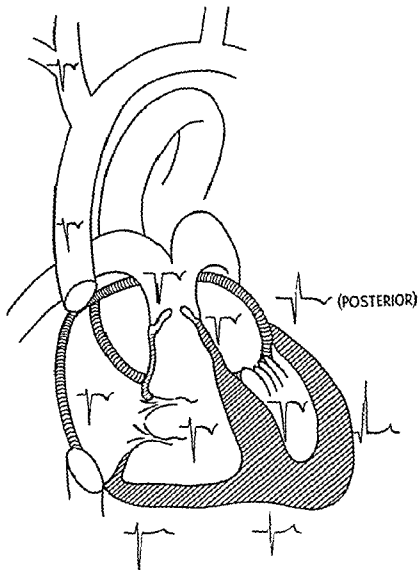


Fig 3 07—The basic QRS T patterns

UNIPOLAR LIMB LEADS

The potential values in the right arm (VR) left arm (VL) and left leg (VF) may be obtained by placing the exploring electrode on the desired limb and linking it with Wilson's neutral electrode As unipolar limb leads are of low voltage it is customary to alter the standardisation so that a

potential difference of 1 millivolt causes a deflection of 15 mm (instead of 10 mm) Alternatively Goldberger's augmented leads may be used. With this technique the V lead is attached to the limb the potential values of which are being measured whilst the wire connecting this limb with the

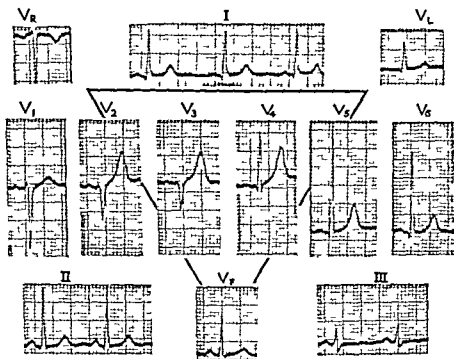


Fig 308 —Unipolar limb leads (V_L V_R V_F) and standard leads 1, 2 and 3
(a) Normal (the heart is more horizontal than vertical)

central neutral terminal is detached and left hanging free. The potentials are thus increased by 50 per cent (Goldberger 1942) thus

$$\text{assuming } V_R + V_L + V_F = 0$$

$$\text{then } V_L + V_F = -V_R$$

Now when an electrode on the right arm is paired with a central terminal linked to the left arm and left leg the galvanometer records $V_R - \frac{V_L + V_F}{2}$

the latter being the mean potentials of the left arm and left leg

$$\begin{aligned} \text{Now } V_R - \frac{V_L + V_F}{2} \\ &= V_R - \frac{(-V_R)}{2} \\ &= V_R + \frac{1}{2}V_R = 1\frac{1}{2}V_R \end{aligned}$$

The reduction in the voltage of R from V₁ to V₃ is due to the resistance of breast tissue and is not uncommon in women

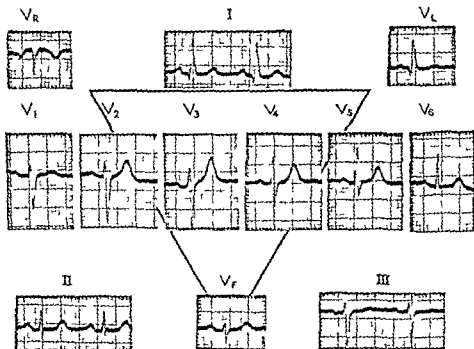


Fig 308 (b) - Horizontal heart

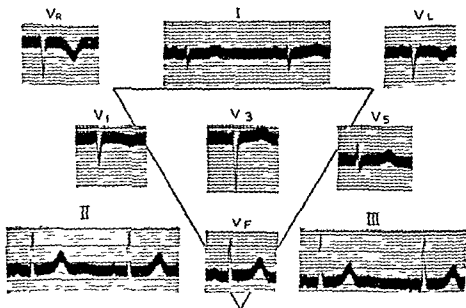


Fig 308 (c) - Vertical heart

Unipolar limb leads are useful in determining the electrical position of the heart in explaining the difference between CR, CF and V chest leads and in demonstrating the basis of the standard leads. VR usually shows inversion of all complexes because it reflects the negative potential of the cardiac cavities transmitted through the great vessels (figs 3.07 and 3.08). When the heart is normal in size and position, VI and VF are mainly positive, dominant left ventricular surface potentials being transmitted more or less equally to both of them, but VF is more positive than VL because the latter is also influenced by the negative potentials of the cavities transmitted through the great vessels (fig 3.08a). When the heart is electrically horizontal, however, left ventricular surface potentials are transmitted more strongly to the left arm and right ventricular surface potentials to the left leg. There is then a small Q and tall R wave in lead VL and a small R and deep S wave in lead VF (fig 3.08b). When the heart is electrically vertical, the negative potentials of the cavities are transmitted more strongly to the left arm and the left ventricular surface potentials more strongly to the left leg. There is then a small R and deep S wave in VL and a small Q and tall R wave in VF (fig 3.08c).

The differences between CR, CF and V chest leads may now be appreciated. CR leads are V leads minus the potentials in VR, whilst CF leads are V leads minus the potentials in VF. As VR potentials are negative, their subtraction from V in CR records makes all deflections more positive – not only is R taller in lead CR_r but T is invariably upright in adult and in children over eight years of age. Again, since VF potentials are normally positive, their subtraction from V in CF records makes all deflections more negative. As the voltage is usually higher in VR than in VF, however, CR leads show greater differences from V leads than do CF leads.

STANDARD LEADS

Einthoven's bipolar leads, introduced at the beginning of the century and adopted as the standard leads throughout the world, consist of the left and right arm (lead 1), the left leg and the right arm (lead 2), and the left leg and left arm (lead 3). Electrocardiograms derived from these leads can be calculated, of course, from the deflections obtained with unipolar limb leads: for lead 1 equals VL–VR, lead 2 equals VF–VR, lead 3 equals VF–VL. The subtraction of the negative potentials in VR from the positive potentials in VL result in strongly positive QRS and T deflections. Again, as the voltage of R in VF is usually higher than that in VL, QRS is also normally positive in lead 3.

By definition there is an obvious relationship between the three standard leads

$$\text{lead 2} = \text{lead 1} + \text{lead 3}$$

This merely states that

$$\begin{aligned}\text{VF}-\text{VR} (\text{lead 2}) &= \text{VL}-\text{VR} (\text{lead 1}) + \text{VF}-\text{VL} (\text{lead 3}) \\ &= \text{VF}-\text{VR}\end{aligned}$$

and has nothing to do with Einthoven's theory or triangle.

The relationship between the standard leads and the Wilson unipolar limb leads is as follows

$$VL = \frac{I+III}{3}$$

$$VR = \frac{I+II}{3}$$

$$VF = \frac{II+III}{3}$$

The augmented values obtained with Goldberger's technique may be derived from the standard leads by changing the denominator in the above equations from 3 to 2

NORMAL APPEARANCES

(Fig 3 08)

P wave P represents the excitation process as it spreads from the sino auricular node through both auricles. It is usually blunt and is upright in leads 1 and 2 but may be inverted in lead 3. Its height should not exceed 2.0 mm and its duration 0.1 second. Following P slight depression of the base line, sometimes hidden by the QRS complex, may be evident and represents auricular recovery or repolarisation. It has been termed the auricular T wave or Ta wave.

P-R interval No deflection is caused by the passage of the excitatory impulse down the bundle of His, its main branches, and Purkinje network so that there is an iso potential interval between auricular and ventricular events, this is the P-R interval and is conveniently measured from the beginning of P to the beginning of QRS. It commonly ranges between 0.12 and 0.20 second but occasionally even in young subjects it may measure 0.21 or 0.22 second without evidence of heart disease or of general ill health.

The P-R interval is little affected by spontaneous variations in heart rate but may be slightly reduced by atropine and slightly lengthened by carotid sinus compression. Vagal tone has a much greater effect on the sinus node than on A-V conduction.

The QRS complex Q, R, and S when all are present form a triphasic complex representing the spread of the accession wave through the ventricles and are convenient symbols for describing the shape of the initial ventricular deflection. Each is applied to a wave so defined by its direction and by its time relationship to the others. Thus any upward deflection is called R or if there are two such R and R₁. A downward deflection is called Q if it precedes R or if it is the only wave present and S if it follows R.

Q rarely measures more than 1 or 2 mm in leads 1 and 2 and is often absent altogether in lead 3, however it may be conspicuous and may

measure up to one third of the amplitude of R. R should exceed 5 mm in height in the most favourable lead unless the spatial vector is unusually postero-anterior. Slight notching or slurring near its base is common and has no significance. Distortion of the apex of R is rare in normal subjects but may be disregarded when unaccompanied by other changes. S is variable and is greatly influenced by axis deviation which will be considered later.

The whole QRS complex should not exceed 0.1 second in duration and rarely exceeds 0.08 second in normal individuals.

RS-T segment This refers to that short segment between the QRS complex and the T wave, i.e. between the end of the excitatory and the beginning of the recovery processes. In some cases this is so short as to represent merely the RS-T junction. Any deviation of the RS-T segment from the isoelectric potential base line should be regarded with suspicion. Slight deviation of the order of 0.5 mm may be within normal limits yet taken in conjunction with other findings may be highly significant.

It is customary to include the proximal portion of the T wave when describing the shape of the RS-T segment, e.g. whether concave, straight or convex. Speaking in this way, a normal RS-T segment curves gently from its point of origin in the direction of the T wave; it is neither straight nor does it deviate in the opposite direction first.

T wave T represents the recovery process or the regression wave (repolarisation) and is known as the second ventricular deflection. It is normally upright in leads I and 2 but may be inverted in lead 3. It should measure at least 2 mm in amplitude in the most favourable lead.

Q-T interval The interval between the beginning of QRS and the end of T represents the total time occupied by ventricular excitation and recovery. It is inversely proportional to the heart rate, ranging between 0.42 second at a speed of 48 per minute and 0.28 second at a speed of 110. The formula of Bazett (1920) is $Q-T = K \sqrt{C}$ where C represents the cycle length. The constant K is variously given as 0.38–0.39 plus or minus 0.04 and is a trifle longer in women than in men and children.

Taran and Szilagyi (1947) have made the sensible suggestion that the Q-T interval should be recorded as corrected for rate, i.e. as Q-T'. This should equal Bazett's constant K, i.e. the actual Q-T interval when the heart rate is 60 per minute or when the cycle length is one second. Q-T' is easily calculated with the aid of a slide rule when the actual Q-T interval and cycle length are known: for Q-T' (or K) = $\frac{Q-T}{\sqrt{C}}$. The Q-T interval is

lengthened by hypocalcaemia (fig. 3.33) and shortened by digitalis. Prolongation of Q-T' provides valuable evidence of active rheumatic carditis (page 271). There is some evidence that Q-T_e is also lengthened by cardiac enlargement from any cause and shortened by cardiac compression as in pericardial effusion (Van Lingen, 1947).

U wave Following T and coinciding with the super-normal recovery

phase a small, rounded positive deflection, the U wave may be seen. Its significance is not fully understood, but it appears to be exaggerated in chest leads taken from the right of the interventricular septum and to be flattened or even inverted in leads taken from the left of the septum when there is left ventricular hypertrophy and the opposite when there is right ventricular hypertrophy. It may also be inverted in left ventricular surface leads during an attack of angina pectoris. It is accentuated by digitalis.

THE CARDIAC VECTOR

Maximum potential differences within the heart at any given moment may be represented in magnitude and direction by a line of appropriate length and spatial direction (drawn from the hypothetical centre of electrical events) which may be called a vector and its direction a spatial axis. Both magnitude and direction of this vector alter from moment to moment during the phases of ventricular excitation and recovery, but may be

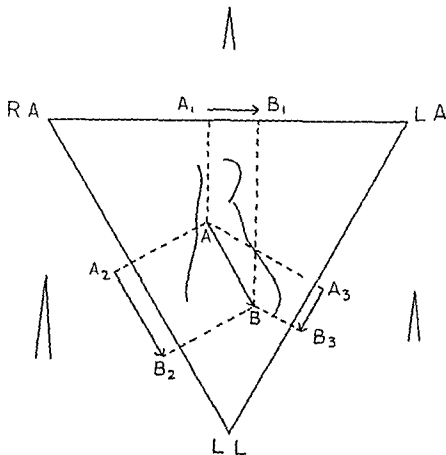


Fig 309—Projection of the frontal plane QRS vector on to the sides of Einthoven's equilateral triangle

resolved into mean values. If such a vector is projected on to the frontal plane of the body, its new momentary or mean manifest value may be calculated by suitable measurements detailed below of the electrocardiograms obtained from any two of Einthoven's leads for the frontal plane or manifest vector may be projected on to the sides of an equilateral triangle the apices of which are represented by the left and right arms (or shoulders) and by the left leg (or symphysis pubis) the sides of the triangle thus representing the three standard leads. For example if the line AB (fig 3 09) represents the maximum momentary manifest QRS vector i.e. if it represents the projection on to the frontal plane of the body of a line in space representing the magnitude and direction of maximum potential differences generated within the heart during the period of ventricular excitation then the lines A₁-B₁, A₂-B₂ and A₃-B₃ obtained by projecting the line AB on to the sides of Einthoven's equilateral triangle give the magnitude and direction of the maximum QRS deflection in leads 1, 2 and 3 respectively. Moreover it can be easily shown that at any given moment the amplitude of the QRS deflection in lead 2 equals the algebraic sum of that in leads 1 and 3 or the amplitude of the QRS deflection in any one lead equals the algebraic sum of that in the other two. The same law applies to auricular activity and to the recovery phase i.e. to the I Ta and T waves and to mean as well as momentary values. Conversely if the magnitude and direction of the QRS complex at any given moment is known in any two leads their resultant drawn from the centre of Einthoven's triangle represents the manifest (frontal plane) vector of QRS at that particular moment and its direction the manifest electrical axis. In current electrocardiographic nomenclature the electrical axis refers to this resultant frontal plane axis as obtained from the maximum upright QRS deflection in any two leads usually R₁ and R₂ if apparently synchronous and is expressed in terms of its angle with the horizontal being plus when rotated clockwise from this base minus when anti clockwise. As so expressed the normal electrical axis lies between 0 and 90 degrees and has a wider range than the frontal plane anatomical axis.

Triaxial reference system. For convenience Einthoven's triangle may be suitably represented as a triaxial reference system (Bayley 1943). The lines representing the three sides of the triangle are transposed so that they intersect at a common point O (fig 3 10). The horizontal line RL then represents lead 1 and the lines RF and LF lead 2 and 3 respectively. The customary signs are preserved so that R is negative, F positive and L negative or positive as shown in the diagram. If the vector OA is projected on to these lines its value in the standard leads may at once be determined by the lengths OA₁, OA₂ and OA₃. The converse may be applied with equal simplicity.

By measuring the net area of QRS in any two leads (instead of momentary synchronous points) by means of a planimeter and suitable magnification (or by dividing the amplitude of the wave by half its width) the area

below the base line being subtracted from that above the resultant mean axis of QRS in the frontal plane can be determined in similar fashion (Wilson *et al* 1934) Measurements may be made in millivolt seconds, microvolt seconds or in suitable units based on voltage \times time (Ashman and Byer 1943) Such a resultant drawn from the centre of Einthoven's triangle having both magnitude and direction is called the mean QRS vector in the frontal plane or the manifest mean QRS vector and its direction the manifest mean QRS axis. Manifest mean vectors for T and P may be similarly obtained. Bayley (1943) has suggested that the symbol

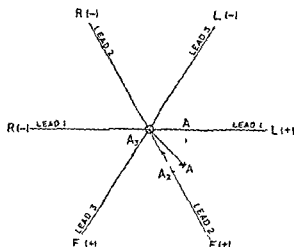


Fig. 3.10—Bayley's triaxial reference system

A might well designate the axis of such vectors, and the symbol A their magnitude. The manifest mean axis of QRS would then be called AQR and its magnitude AQR .

If the heart were a simple uniform muscle block the algebraic net area occupied by QRS and T would be zero as it is not. The net area of QRST has a positive or negative value which if measured in any two leads may be resolved into a vector drawn from the centre of Einthoven's triangle. The axis of this vector or the manifest mean QRST axis (\bar{A}_{QRST}) has been called the ventricular gradient (Wilson, Macleod and Barker 1931) or G and its magnitude G . The vector represents the magnitude and direction of maximum local variations in the speed of the processes of excitation and recovery whereby the heart differs from a uniform muscle block.

The manifest mean axis of QRS averages about 60 degrees that of T about 50 degrees. The ventricular gradient in hearts which are not anatomically rotated ranges between 45 and 65 degrees. On the whole, hearts which are relatively central in position i.e. rotated clockwise (viewed from the front) about their antero-posterior anatomical axis are also rotated clockwise (viewed from the apex) about their longitudinal anatomical axis and show clockwise deviation i.e. deviation to the right of all manifest momen

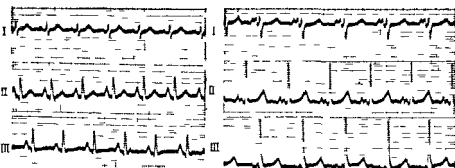
tary and mean electrical axes but the greatest shift occurs with the ordinary momentary electrical axis of QRS and the least with the ventricular gradient. This also applies to transverse hearts with anti clockwise rotation and deviation of all electrical axes to the left (Ashman and Byer 1943).

From what has been said it should be clear that the QRS and T vectors in the frontal plane of the body alter in magnitude and direction from moment to moment during the phases of ventricular excitation and recovery respectively. As one end of such a vector is fixed at the centre of Einthoven's triangle it follows that the other end must describe a continuous curve. Mann (1930) showed how such curves could be reconstructed and later devised a method of recording them directly (1931). More recently Wilson and Johnston (1938) employing the cathode ray oscillograph published typical curves and called them vectorcardiograms. Even these however are restricted to the behaviour of the vector in the frontal plane of the body being so limited by use of the standard limb leads.

ELECTROCARDIOGRAPHIC ABNORMALITIES

ABNORMALITIES OF THE P WAVE

There are four main varieties of P wave deformity: the tall sharp P wave of right auricular hypertrophy (fig 3 11a) the conspicuous widened P wave of left auricular hypertrophy which may be bifid, rounded or flat topped



(a) Right auricular hypertrophy

(b) Left auricular hypertrophy

Fig 3 11—Abnormal P waves

(fig 3 11b) the low voltage widened P wave which may be also bifid, rounded or flat topped (fig 3 11c) and the inverted P wave (fig 3 11d).

Tall sharp P waves are characteristic of tricuspid stenosis, chronic pulmonary heart disease and congenital pulmonary stenosis. The voltage ranges between 2 and 5 mm and as the wave is not widened it becomes peculiarly sharp like an arrowhead. Similar P waves are sometimes seen in mitral stenosis, thyrotoxicosis, massive pulmonary embolism, asthma and pneumonia. They are usually most evident in leads 2 and 3.

Conspicuous widened P waves measuring 0.12 second in duration are almost diagnostic of mitral stenosis. The voltage usually approaches 2.5 mm but rarely exceeds it. Many examples are bifid but others are blunt or flat topped. They are usually seen best in leads I, 2 and V₅.

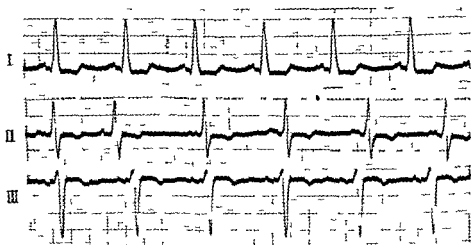


Fig. 3 11 (c)—Hypertensive low voltage

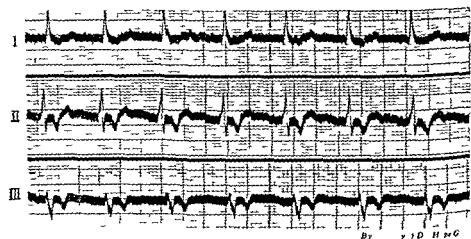


Fig. 3 11 (d)—Inverted (nodal)

P waves similar in shape and width but usually of lower voltage may be seen sometimes in advanced cases of hypertensive heart disease or of aortic valve disease. It is uncertain whether they represent left auricular dilatation due to left ventricular failure as originally suggested by Wood and Selzer (1939) or inter atrial block (Berconsky and Klotzman 1945).

Inverted P waves are found in lead I in cases of dextrocardia and in all leads in many cases of nodal rhythm. Occasionally P is inverted in leads 2 and 3 without obvious cause.

ABNORMALITIES OF THE QRS COMPLEX

Axis deviation It has already been pointed out (page 81) that the electrical axis of the heart refers to the frontal plane projection of the maximum momentary spatial vector and usually lies between 0 and 90 degrees more or less in the anatomical axis. Anti clockwise rotation of the

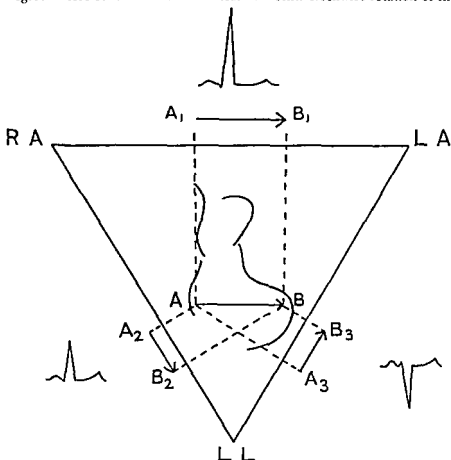


Fig 3 12—Left axis deviation (F nthoven's triangle)

heart about its antero posterior axis (viewed from the front) or about its longitudinal axis (viewed from the cardiac apex) causes deviation of the electrical axis to the left so that the frontal plane vector may make a minus angle with the horizontal whilst clockwise rotation about similar axes causes right axis deviation the vector now making an angle of more than 90 degrees with the horizontal. Left or right axis deviation respectively also occurs when the left or right ventricle is disproportionately enlarged. Moreover left ventricular enlargement is often associated with anti clockwise rotation about both anatomical axes and right ventricular enlargement with clockwise rotation.

Reference to Einthoven's triangle will show that if the electrical axis deviates to the left and approaches or surpasses the horizontal lead 1 becomes the axial lead (fig 3 12) R_1 then carries the maximum voltage R_2 is smaller and the maximum QRS deflection in lead 3 is downwards i.e. the main deflection is S. In such cases S is really the electrical counterpart of R_1 . Unipolar limb leads commonly show an electrically horizontal heart R in VL and S in VF being unusually conspicuous.

Left axis deviation occurs in 10 per cent of normal individuals in any condition in which the left ventricle is disproportionately enlarged in cardiac displacement to the left from scoliosis or from intrathoracic causes and when the diaphragm is elevated causing the heart to lie more transversely. It may not be possible from examination of the limb lead QRS complexes alone to decide whether axis deviation is due to displacement or to left ventricular preponderance but this distinction may often be made by considering the behaviour of the RS-T segment and T wave

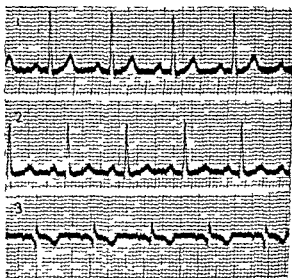


Fig 3 13—Axis deviation due to elevation of the diaphragm ($Q_3 S_1$ type)

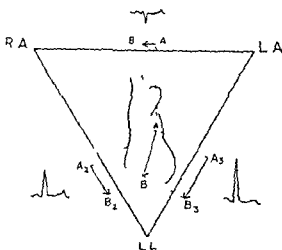


Fig 3 14—Right axis deviation (Einthoven's triangle)

and especially by noting the QRS pattern in multiple chest leads (*vide infra*)

A particular form of axis deviation is seen with elevation of the diaphragm as from obesity, pregnancy, flatulence or ascites R_1 is taller than R_2 , S_1 and Q_3 are prominent and T_3 is inverted (fig 3 13). In such cases there is no Q wave in lead VF and the T wave usually remains inverted in lead V1.

When the electrical axis is deviated to the right so that it occupies a more or less vertical position lead 3 becomes the axial lead (fig 3 14) R_3 then carries the maximum voltage R is smaller whilst the maximum deflection in lead 1 is S which is the electrical counterpart of R_3 . In unipolar limb leads S is conspicuous in V_1 and R in V_F . Right axis deviation is the rule in newly born infants is common in very young children occurs in 1 per cent of normal children over the age of eight and is rarely seen in

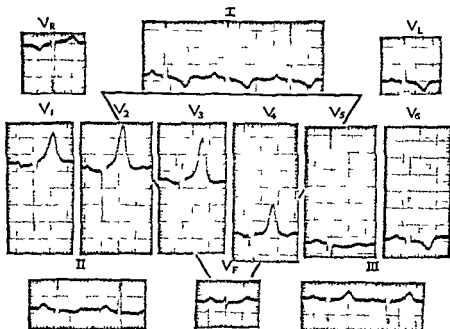


Fig 3 15—Left ventricular preponderance (heart horizontal)

strictly normal adults. It may be caused by appropriate cardiac displacement or rotation and by right ventricular dominance. As with left axis deviation it may not be possible from inspection of the limb lead QRS complexes alone to determine whether the axis shift is due to right ventricular dominance or otherwise, but the behaviour of QRS in multiple chest leads may clarify the issue (*vide infra*).

Left ventricular preponderance. When the left ventricle is enlarged the accession wave takes longer to penetrate that chamber and creates more powerful potential differences. Thus R in leads V_5 and V_6 and S in leads V_1 and V_2 have a larger amplitude (R in V_4 5 or 6 $>$ 25 mm, S in V_1 $>$ 15 mm). The intrinsic deflection in left ventricular surface leads is delayed (longer than 0.05 second) and the width of QRS slightly increased (0.1 to 0.12 second). In addition the transition zone is usually shifted to the left the heart being horizontal and R is found to be exceptionally small in leads V_1 and V_2 . Secondary changes in the T wave occur in advanced cases the

R-T segment being depressed and T inverted in leads V_5 and V_6 and the S-T segment being elevated and T sharply upright in leads V_1 and V_2 (fig 3 15)

When the heart is horizontal which is usual V_L resembles V_5 and V_F resembles V_1 both in respect of QRS and T. The appearances in standard lead 1 therefore also resemble V_5 or V_6 and those in lead 3 resemble V_1

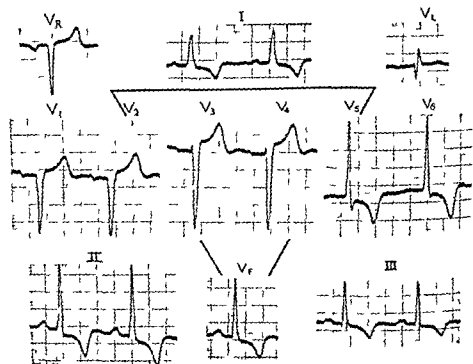


Fig 3 16—Left ventricular preponderance (heart semi vertical)

When the heart is more or less vertical which is less common left ventricular surface potentials are transmitted more to the left leg. There is then no axis deviation in standard leads (Wilson 1944) but high voltage and perhaps T wave inversion in all (fig 3 16). Concordant left ventricular preponderance as it is called is best seen in concentric left ventricular hypertrophy such as may occur in aortic stenosis and malignant hypertension.

Right ventricular dominance When there is gross enlargement of the right ventricle the potential differences generated by the wall of that chamber may approach or even surpass those from the left ventricle. Right ventricular surface leads may then truly represent the outward spread of the accession wave beneath the exploring electrode. R is therefore taller than usual in V_1 and may be the chief ventricular deflection at the same time the intrinsic deflection is delayed and S is conspicuous in V_5 and V_6 (fig

hypertrophy is usually associated with high voltage whereas in bundle branch block QRS is commonly notched splintered or heavily slurred. When the heart is grossly dilated there may be some delay in the passage of the excitatory impulse down the Purkinje network causing intraventricular block. Some such mechanism may account for the transient right bundle branch block that occurs occasionally in massive pulmonary embolism and for the right bundle branch block 'so commonly seen with atrial septal defect. A Q wave can nearly always be demonstrated in suitable left ventricular surface leads when widening of the initial ventricular deflection is due to left ventricular hypertrophy, whereas it is ordinarily absent in left bundle branch block.

Widening of the QRS complex is also seen in uræmia when it is due to a raised blood potassium (figs 3.33 and 3.34).

Bundle branch block In *left bundle branch block* the excitatory process spreads through the right ventricle in normal fashion but does not at first reach the left ventricle. As the interventricular septum is excited from the right side the accession wave spreads through it from right to left. The cavity of the left ventricle therefore becomes initially positive and this potential is transmitted passively to the surface as an R wave in V₅ or V₆. There can be no Q wave in such leads with a healthy septum. When the accession wave reaches the left side of the septum there is an immediate reversal of polarity the left ventricular cavity becoming momentarily negative. This negativity is again transmitted passively to the surface V₅ showing a momentary downward deflection following the initial R wave. Almost immediately however the excitatory process spreads throughout the endocardium of the left ventricle and the accession wave begins to flow outwards in the usual way. The surface of the left ventricle then becomes actively positive and the true R wave is written. When the surface is activated the final intrinsic downward deflection occurs. V₅ or V₆ thus exhibits a large widened R wave interrupted by a relatively early notch representing the arrival of the accession wave at the left side of the septum (fig 3.18). Right ventricular surface potentials are influenced at first by a normal right ventricular accession wave and later by the delayed negativity of the cavity of the left ventricle which is passively transmitted through the depolarised septum and right ventricle. Thus V₁-V₃ exhibit small R waves early intrinsic deflections and deep wide S waves. The total duration of QRS commonly measures 0.12 to 0.16 second. As the heart is usually horizontal the V₅-V₆ pattern is seen also in VL and lead I and the V₁ pattern in VF and lead 3. Should the heart be more or less vertical however the V₅-V₆ pattern is transmitted to the left leg and QRS may be mainly upright in all standard leads as in concordant left ventricular preponderance. The T wave is secondarily deformed and the RS-T segment deviated from the base line in all leads and are commonly of opposite sign to the QRS complex. Thus with horizontal hearts the RS-T segment is depressed and the T wave inverted in V₅ V₆ VL and standard lead I.

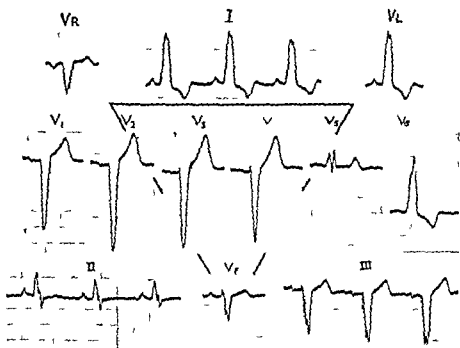
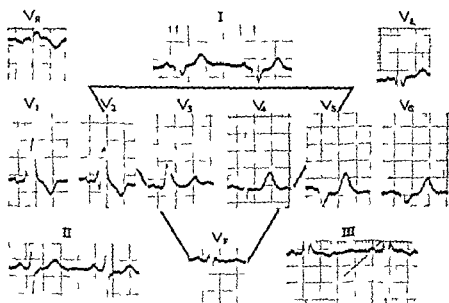
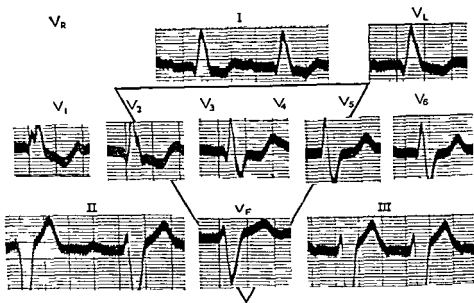


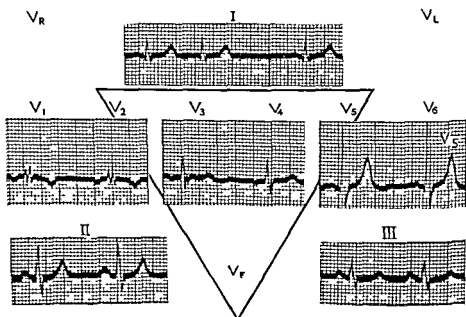
Fig 3 18—Left bundle branch block (heart horizontal)



3 19—Right bundle branch block (usual appearances)



(a) Associated with a vertical heart



(b) Partial seen clearly only in chest leads

Fig 3 20—Right bundle branch block

In *right bundle branch block* the septum is activated entirely from the left side. The potential of the right ventricular cavity is therefore initially positive and is passively transmitted to the surface where it may be recorded as the first part of R. When the accession wave reaches the right side of the septum the polarity is abruptly reversed and a pseudo intrinsic deflection is recorded at the surface. Almost at once however the right ventricular wall is invaded and the surface then becomes actively positive. This results in a second R wave and finally in the true intrinsic deflection. Leads V₁ and V₂ therefore show a widened notched R wave or a large M complex. T is in the opposite direction (fig. 3 19). Over the left ventricle in leads V₅ and V₆ normal QR wave and intrinsic deflection are followed by a grossly slurred S wave representing delayed negativity of the right ventricular cavity passively transmitted through the depolarised septum and left ventricle. As a rule V₅ and V₆ potentials are transmitted to V₁ and form the pattern of standard lead 1; the M complex of V₁-V₂ is usually seen in V₇ and in standard lead 3. When the heart is vertical however V₁ potentials may be transmitted to V₁ and standard leads may look like left bundle branch block (fig. 3 20a). Multiple chest leads may be necessary not only to determine which bundle branch is blocked but also to detect the lesion at all in some cases partial right bundle branch block for instance is frequently overlooked in standard leads (fig. 3 20b).

ABNORMALITIES OF THE RS T SEGMENT AND T WAVE

It is profitable to consider the RS T segment and T wave together and in many cases to consider them also in relationship to the QRS complex for they are all ventricular events. The various patterns made up by these three variables in limb and multiple chest leads provide a wealth of information concerning the state of the ventricles in health and disease. Secondary inversion of the T wave in relation to QRS changes has already been described.

Myocardial infarction It is customary to describe two types of electrocardiogram associated with myocardial infarction. T₁ and T₂ types (Parkinson and Bedford 1927) the first denoting anterior the second posterior lesions (Barnes and Whitten 1949). There is no essential difference in the shape of these two patterns the difference depending upon the leads in which they are found.

If an infarct involves the whole thickness of the muscle wall no accession wave can flow through it. The negative cavity potential produced by outward spread of the accession wave through remote healthy muscle is then passively transmitted through the infarct to the surface overlying it. An electrode placed over the infarct therefore registers an initial negative deflection or Q wave. If the infarct involves only the outer part of the muscle the accession wave begins as usual the surface potential is initially positive and there can be no pathological Q wave but when the accession wave reaches the infarct it can advance no farther so R is reduced in

amplitude. In anterior left ventricular infarcts these QRS changes may be registered in leads V_3 , V_4 , V_5 and V_6 being more marked in V_3 - V_4 in antero septal infarcts and in V_5 - V_6 in antero lateral infarcts. They are commonly transmitted to V_L and are therefore seen well in standard lead 1 (fig 3 21). Similar QRS changes occur in posterior infarcts but are transmitted to V_F and thus to standard lead 3 (fig 3 22a). When the heart is vertical however typical changes in V_5 from an anterior infarct may be transmitted to lead V_F and hence to standard leads 2 and 3 (fig 3 22b).

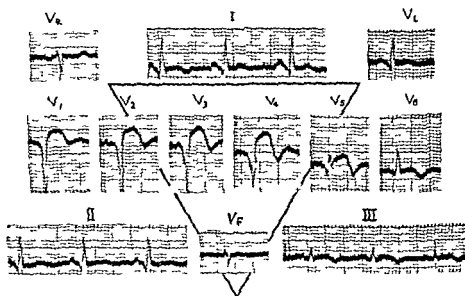


Fig 3 21.—Anterior myocardial infarction showing pathological Q waves and elevation of the RS-T segment in leads V_1 - V_6 , V_L and standard limb lead 1

Partly necrosed muscle sets up a steady current due to the development of potential differences between injured and healthy tissue. Injured tissue is electro negative, healthy tissue is positive and completely necrosed tissue electrically inert. When the injured area involves the outer portion of the ventricular wall the surface is therefore negative, the current flowing from without inwards (Wilson *et al.* 1933). An electrode placed over the infarct registers this negativity by depressing the base line. This is shown in the electrocardiogram (fig 3 21) by abrupt elevation of the base line when the current of injury is momentarily abolished by spread of the accession wave through the healthy tissue, for such activation causes the healthy tissue to take up a negative potential and so abolishes the potential differences set up by the injury. In other words, a recent deep infarct associated with superficial injury results in elevation of the RS-T segment. In anterior infarcts this displacement is seen in leads V_3 - V_6 and is commonly transmitted to V_L and hence to standard lead 1 (fig 3 21). In posterior infarcts

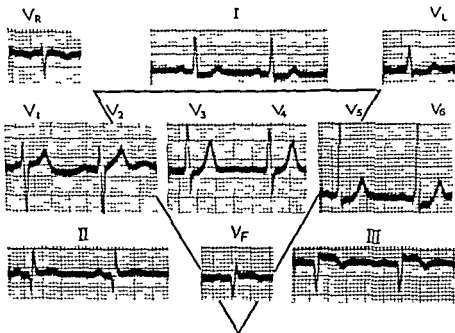


Fig 3 22 (a)—Posterior myocardial infarction showing pathological Q waves and elevation of the RS-T segment in lead V_F and standard limb lead 3

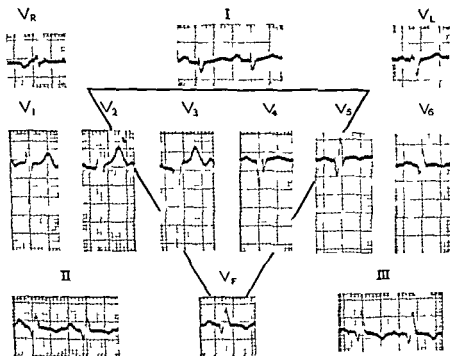
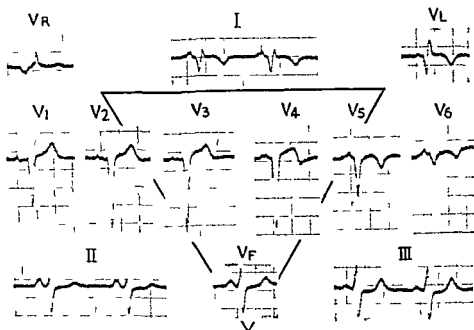
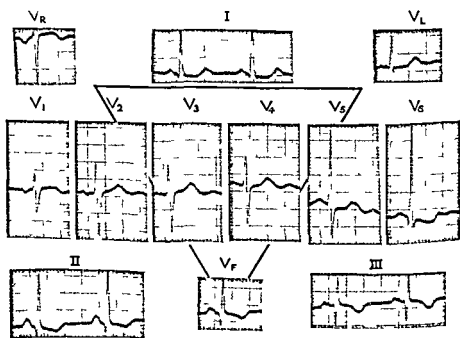


Fig 3 22 (b)—Anterior myocardial infarction with vertical heart. Standard leads show changes that simulate those of posterior infarction



(a) Anterior



(b) Posterior

Fig 3 23—Later stages of anterior (a) and posterior (b) infarction showing typical Q waves and inversion of the T wave in appropriate leads

it is seen in low α sphigeal leads in V_7 and is transmitted to V_F and hence to standard lead 3 (fig 3 22a)

Pathological Q waves may be seen in acute cases within a few hours of the onset and usually outlast all other evidence of infarction often being permanent Elevation of the RS T segment also occurs early but usually subsides within two or three weeks The shape of the segment is typical,

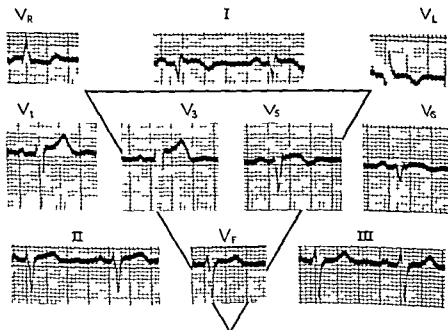
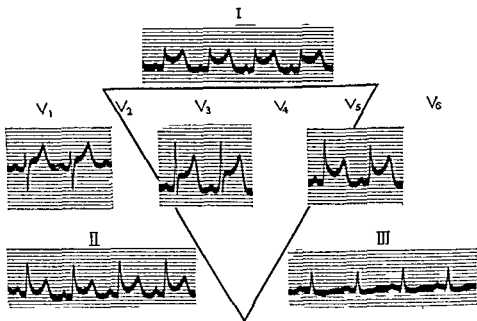


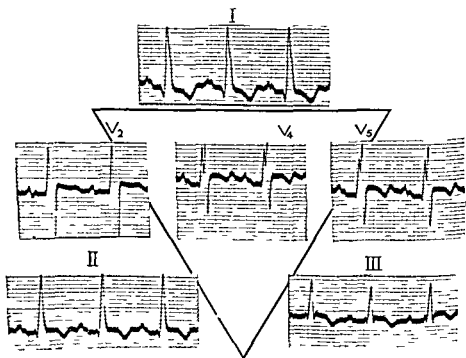
Fig 3 24—Anterior myocardial infarction showing an R wave which is smaller in V_3 to V_6 than in V_1

being straight instead of concave when elevated and being convex or cove shaped (Pardee 1920) when the RS T junction approaches or regains the iso potential level The T wave itself becomes inverted within a few days of the onset often profoundly so reaching its greatest amplitude at about the same time that the RS T junction first regains the iso potential level (fig 3 23 a and b) Further changes are regressive but the appearances rarely revert to normal

In T_1 patterns reciprocal effects are usually observed in lead 3 i e the RS T segment may be depressed at first and T may be sharply upright later Again in posterior infarcts early RS T depression and later accentuation of the T wave may often be seen in lead 1 and in anterior chest leads A helpful sign of old anterior infarction is an R wave in V_1 - V_2 which is taller than that in V_3 - V_4 (fig 3 24) especially when the appearances in V_5 - V_6 are more or less normal Finally it is most important to understand that characteristic changes may be found in multiple chest



(a)—Early stage showing elevation of the RS T segment



(b)—Late stage showing inversion of the T wave in all leads

Fig. 325—Pericarditis

leads or in an oesophageal lead when the standard limb leads are normal and that a single chest lead may be normal when others show diagnostic features

Pericarditis In all types of generalised pericardial disease except hydropericardium superficial epicardial involvement may cause a current of injury to flow from the surface towards the underlying healthy muscle in other words the surface of the heart develops a negative potential. The

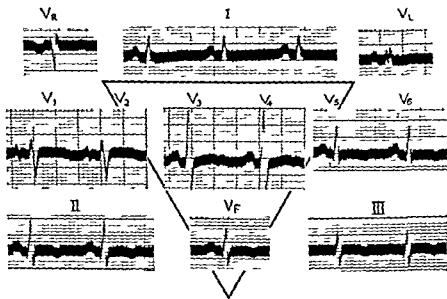


Fig. 3.26—Chronic constrictive pericarditis showing low voltage and flat T waves.

situation therefore resembles that in myocardial infarction but the lesion is general instead of local. Thus in the initial stages elevation of the RS-T segment may be seen in all chest leads in both V₁ and V₆ and therefore in all standard leads (fig. 3.25a). Unlike most records of acute myocardial infarction the RS-T segment remains concave. As the underlying muscle is healthy there are no pathological Q waves. After a few days the RS-T segment regains the isopotential level and the T wave becomes inverted (fig. 3.25b). Upward coving of the RS-T segment does not occur. If pericarditis is localised the changes described may be confined to corresponding leads but few important forms of pericarditis remain localised for long. Serial records nearly always reveal what may be called the T pattern in contrast to the T₁ or T₂ types of myocardial infarction. Low voltage QRS complexes usually indicate pericardial effusion.

The electrocardiogram returns to normal as the pericarditis recovers. In chronic constrictive pericarditis flattened or inverted T waves in all leads are permanent and are usually associated with low voltage QRS complexes.

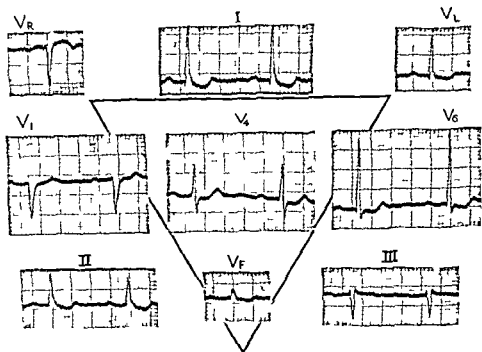


Fig 3 27 (a)—The effect of digitalis on the electrocardiogram showing sagging of the RS T segment and shortening of Q Tc to 0.33 second

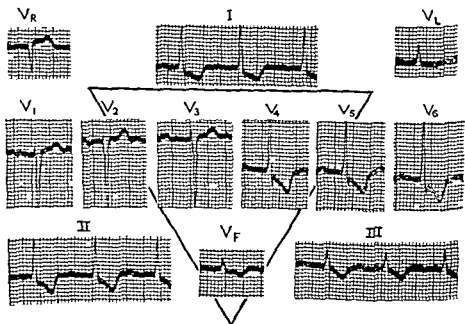


Fig 3 27 (b)—Showing gross depression of the RS T segment or an inverted T wave with a straight proximal limb Q Tc is shortened to 0.36 second

(fig 3 26) Not infrequently the P waves are widened and relatively prominent

Digitalis T wave pattern Digitalis depresses the RS T segment and shortens the QT interval. At first the RS T junction is depressed and there is downward curving of the RS T segment T remaining upright (fig 3 27a). In the second stage sagging is more marked and the peak of T can no longer be discerned. In extreme digitalisation the RS T segment becomes a straight line sloping downwards from its depressed origin to a blunt peak (fig 3 27b).

In normal hearts these effects are seen in all leads but especially in lead V₅ and standard lead V₂. When the heart is electrically horizontal they are seen best in V₅ V₆ and in standard lead 1 when it is electrically vertical they are best seen in V₅ V₆ and in standard lead 3. When the left ventricle is enlarged and the heart is horizontal the changes occur more markedly in V₅ V₆ and standard lead 1 and the RS T segment may be elevated and upwardly convex in V₁ V₆ and standard lead 3. When the right ventricle is enlarged they may be most conspicuous in V₁ V₆ and standard lead 3 and the RS T segment may be elevated and upwardly convex in V₅ V₆ and standard lead 1.

Anoxic T waves Electrocardiograms taken from patients during an attack of angina pectoris may show transient depression of the RS T

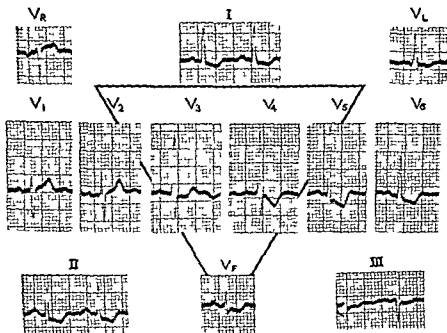


Fig 3 28 (a)—Depression of the RS T segment during an attack of angina pectoris

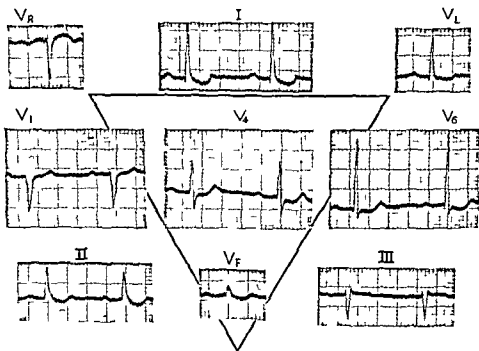


Fig 3 27 (a)—The effect of digitalis on the electrocardiogram showing sagging of the RS-T segment and shortening of Q-Tc to 0.33 second

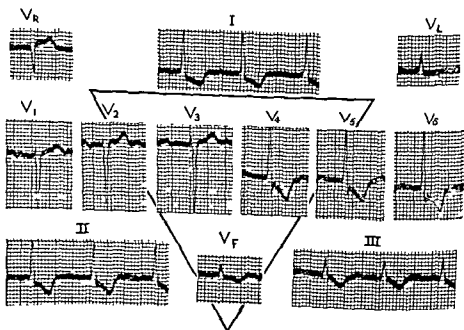


Fig 3 27 (b)—Showing gross depression of the RS-T segment or an inverted T wave with a straight proximal limb Q-Tc is shortened to 0.36 second

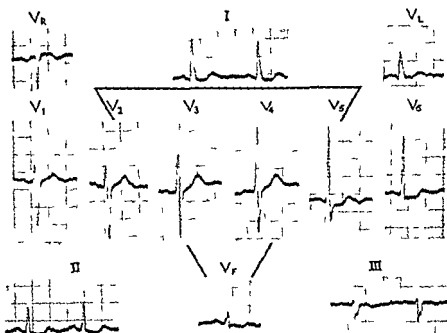


Fig. 330—Relatively permanent depression of the RS-T segment in left-ventricular surface leads in a case of angina pectoris due to occluded disease of the coronary arteries.

Myocardema pattern. Flat or inverted T waves in all leads are characteristic of myocardema (fig. 331). In such cases the voltage of QRS is usually below 6 millimetres in the most favourable standard lead, and there is often bradycardia. Similar appearances may be found in chronic constrictive pericarditis in long-standing cases of severe anaemia, particularly pernicious, and in anoxic chronic pulmonary heart disease, but in these there is commonly tachycardia. In severe cases of ischaemic heart disease with

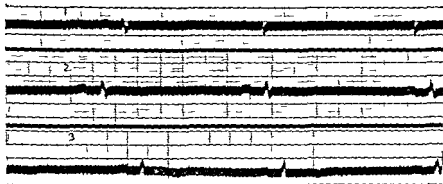


Fig. 331—Myocardema.

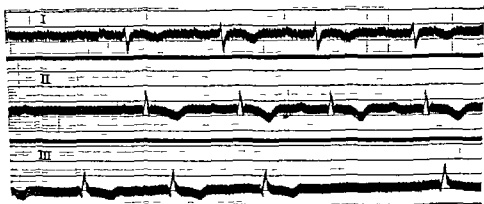


Fig 332—Pneumonic carditis. There is partial heart block with dropped beats and inversion of the T wave in all leads

repeated myocardial infarction somewhat similar graphs may be encountered. Indeed, when the whole heart is involved in any disease and when recurrent heart failure has occurred, the voltage of QRS may be low and the T waves flat or slightly inverted in all leads whatever the etiology.

Carditis pattern. In any form of carditis, but especially in diphtheria and least frequently in acute rheumatism, simple inversion of the T waves may occur and may favour any lead (fig 332). The RS-T segment may be normal or depressed. The voltage of QRS is usually normal.

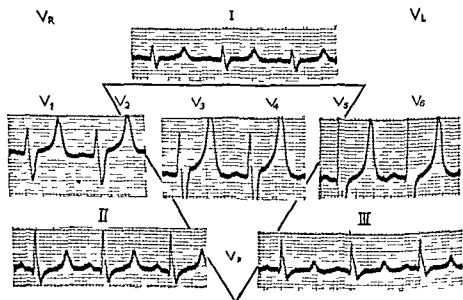


Fig 333—High voltage, sharply peaked T waves in uræmia associated with a high blood potassium. The long QT interval is due to hypocalcæmia. Widening of QRS due to potassium is well seen in the chest leads.

Potassium T waves In uræmia when the blood potassium is high unusually sharp T waves of high voltage are often seen (fig 3 33). Similar T waves may be produced in normal subjects by raising the blood potassium to about 25 mg per cent by giving 10 to 20 G of potassium acetate by

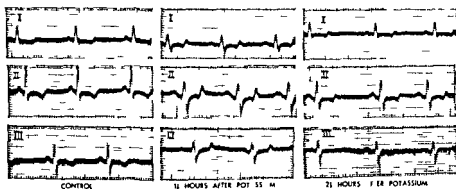


FIG 3 34—Effect of potassium on the T waves in a case of concordant left ventricular preponderance. The QRS complex is also widened.

mouth.* A high blood potassium also tends to rectify many forms of inverted T wave (fig 3 34) but not those due to myocardial infarction which may be exaggerated (Sharpey Schafer 1943). Widening of P and QRS is also due to potassium and is seen in both illustrations.

When the blood potassium is unduly low (<12 mg per cent) the S-T segment and T wave may be depressed and the P-R interval and Q-T_c prolonged (Perelson and Cosby 1949).

REFERENCES

- Ashman R (1948) The physiological and physical aspects of the electrocardiogram. *The Chest and the Heart*, ed by Myers J A and McKinlay C N. Vol II, p 1421.
- and Byer E (1943) The normal human ventricular gradient I. Factors which affect its direction and its relation to the mean QRS axis. *Amer Heart J* 25: 16.
- (1943) The normal human ventricular gradient II. Factors which affect its manifest area and its relationship to the manifest area of the QRS complex. *Ibid* 25: 36.
- Barnes A R and Whitten M B (1929) A study of the R-T interval in myocardial infarction. *Ibid* 5: 142.
- Bayley R H (1943) On certain applications of modern electrocardiographic theory to the interpretation of electrocardiograms which indicate myocardial disease. *Ibid* 26: 769.
- Bazett H C (1920) An analysis of the time relations of the electrocardiogram. *Heart* 7: 353.
- Bell G H, Knox J A C and Small A J (1939) Electrocardiograph electrolytes. *Brit Heart J* 1: 229.

Berconsky I and Klotzman M (1945) Significado de ciertas alteraciones de la onda P del electrocardiogram P de bajo voltaje ancha y bifida *Medicina* 5 347

Craib W H (1930) The electrocardiogram MRC Special Report Series No 147 London

Curtis H J and Cole K S (1941) Membrane resting and action potentials of the squid giant axon *Amer J Physiol* 133 254

Einthoven W (1903) Die galvanometrische Registrierung des menschlichen elektrokardiogramms zugleich eine Beurtheilung der Anwendung des Capillar elektrometers in der Physiologie *Pflüger's Arch f d ges Physiol* 99 472

Goldberger E (1942) A simple indifferent electrocardiographic electrode of zero potential and a technique of obtaining augmented unipolar extremity leads *Amer Heart J* 23 483 — (1947) Unipolar lead electrocardiography London

Jenks J L and Graybiel A (1935) Electrode Jelly *Amer Heart J* 10 693

Kolliker A and Müller H (1856) Nachweis der negativen Schwankung des muskelstroms am natürlich sich contrahirenden muskel *Verhandl d phys med Gesell z Würzburg* 6 528

Levy R L Barach A L and Bruenn H G (1938) Effects of induced oxygen want in patients with cardiac pain *Amer Heart J* 15 187

Lewis T (1925) The mechanism and graphic registration of the heart beat London 3rd ed

Mann H (1900) Method of analyzing the electrocardiogram *Arch intern Med* 25 483 — (1931) Interpretation of bundle branch block by means of monocardigram *Amer Heart J* 6 447

Pardee H E B (1900) An electrocardiographic sign of coronary artery obstruction *Arch intern Med* 26 244

Parkinson J and Bedford D E (1927) Successive changes in the electrocardiogram after cardiac infarction (coronary thrombosis) *Heart* 14 195

Perelson H N and Cosby R S (1949) The electrocardiogram in familial periodic paralysis *Amer Heart J* 37 1126

Præcordial Leads in Electrocardiography (1938) A joint memorandum of a Committee of the Cardiac Society of Gt Britain and Ireland and the Committee of the Amer Heart Ass *Brit med J* 1 187

Robb J S and Robb R C (1938) Abnormal distribution of the superficial muscle bundles in the human heart *Amer Heart J* 15 597

Schlamowitz I (1946) An analysis of the time relationship within the cardiac cycle in electrocardiograms of normal man *Ibid* 31 329

Sharpey Schafer E P (1943) Potassium effects on T wave inversion in myocardial infarction and preponderance of a ventricle *Brit Heart J* 5 80

Standardisation of Præcordial Leads Supplementary Report by the Committee of the Amer Heart Ass (1938) *Amer Heart J* 15 235

Taran L M and Szilagyi N (1947) The duration of the electrical systole (QT) in acute rheumatic carditis in children *Amer Heart J* 33 14

Van Lingen B (1947) Electrocardiographic radiological venous pressure circulation time and exercise tolerance test studies in the diagnosis of heart disease being a thesis submitted for the degree of Doctor of Medicine of the University of Witwatersrand Johannesburg

Waller A D (1887) A demonstration on man of electromotive changes accompanying the heart's beat *J Physiol* 8 209

Wilson F N (1930) The distribution of the potential differences produced by the heart beat within the body and at its surface *Amer Heart J* 5 599

- Johnston I D (1938) The vectorcardiogram *Ibid* 16 14
- — Macleod A G and Barker P S (1934) Electrocardiograms that represent the potential variations of a single electrode *Ibid* 9 447
- — Rosenbaum F I and Barker P S (1946) On Einthoven's triangle the theory of unipolar electrocardiographic leads and the interpretation of the precordial electrocardiogram *Ibid* 32 277
- Macleod A G and Barker I S (1931) The interpretation of the initial deflections of the ventricular complex of the electrocardiogram *Ibid* 6 637
- — (1931) The T deflection of the electrocardiogram *Tr Am Physicians* 46 29
- — — (1933) The distribution of the currents of action and of injury displayed by heart muscle and other excitable tissues Ann Arbor
- — — Johnston F G (1934) The determination and the significance of the areas of the ventricular deflections of the electrocardiogram *Amer Heart J* 10 46
- *et al* (1944) The precordial electrocardiogram *Ibid* 27 19
- Wolferth C C and Wood F C (1932) The electrocardiographic diagnosis of coronary occlusion by the use of chest leads *Amer J med Sci* 183 30
- — (1932) Further observations upon the use of chest leads in the electrocardiographic study of coronary occlusion *Al Clin North Amica* 16 161
- — (1932) An electrocardiographic study of experimental coronary occlusion the inadequacy of the three conventional leads in recording certain characteristic changes in action current *J clin Invest* 11 815
- — (1933) Experimental coronary occlusion *Arch intern Med* 51 771
- Wood P H and Selzer A (1939) Chest leads in clinical electrocardiography *Brit Heart J* 1 49
- — (1939) A new sign of left ventricular failure *Ibid* 1 81

CHAPTER IV

DISORDERS OF CARDIAC RHYTHM

THE speed and regularity of the heart beat are controlled by the sino auricular node of Keith and Flack (1907) situated in the upper part of the sulcus terminalis anterior to and to the right of the mouth of the superior vena cava (fig 4 01). Approximately 70 times per minute this node discharges itself and initiates an excitation wave which spreads in all directions over both auricles. Close to the opening of the coronary sinus above the base of the tricuspid valve on the right side of the atrial

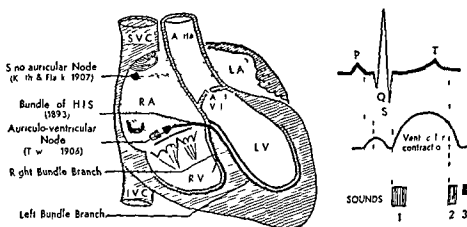


Fig 4 01—Anatomy of the conducting system

septum is situated the auriculo ventricular node of Tawara (1906). This also forms impulses but at a slower rate so that normally it is prematurely discharged by the excitation wave initiated by the S A node. The impulse then spreads down the Bundle of His which passes horizontally to the left to penetrate the membranous interventricular septum where it divides into left and right bundle branches. These pass down each side of the muscular septum just beneath the endocardium. The bundle branches then break up into a network of Purkinje fibres which carry the excitatory process to the sub endocardial myocardium.

Physiology of conduction From the 'pace maker' in the sino auricular node the excitation wave spreads through auricular muscle at a speed of about 1 000 mm per second. Passage through the A V nodal tissue is believed to be relatively slow and is estimated at 200 mm per second. Spread down the bundle branches and the Purkinje fibres is rapid and is

probably as fast as 400 mm per second. Conduction through the ventricles which is believed to proceed directly outwards is put at 400 mm per second (Lewis 1925).

Both the S A and A V nodes are under direct autonomic control, being stimulated by sympathetic activity and depressed by vagal activity. Cardiac accelerator nerves arise from the lateral horns of the upper 4th or 5th dorsal segments of the spinal cord, enter the sympathetic chain and pass cranially to the cervical ganglia. Post ganglionic fibres form the superior, middle and inferior cardiac nerves which terminate in the S A and A V nodes.

IRREGULARITIES AND ALTERATION OF HEART RATE INITIATED OR GOVERNED BY THE SINO AURICULAR NODE

SINUS ARRHYTHMIA

There is probably no such thing as an absolutely regular heart. Slight irregularity, the heart quickening with inspiration and slowing with expiration, is normal and depends upon variations in vagal tone governed by a reflex which is thought to be initiated by receptors in the lungs. Another

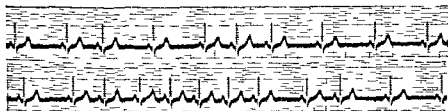


Fig. 402—Sinus arrhythmia

form of sinus arrhythmia occurs independently of respiration. Both are more common in the young and when the heart rate is slow, tend to be exaggerated by drugs which increase vagal tone (such as digitalis) and may be abolished by exercise or by atropine.

Other varieties of sinus arrhythmia are not essentially different but owe their recognition to some particular associated feature: thus there is a form associated with sino auricular block, another with sinus bradycardia and paroxysmal auricular fibrillation or flutter, a third with convalescence from certain infectious fevers, especially influenza, and so on. Increased vagal tone is common to all these types.

Diagnosis is usually easy, or doubt is soon resolved by means of exercise, atropine, or amyl nitrite. An electrocardiogram provides conclusive evidence (fig. 402).

Although sinus arrhythmia is normal, it should not be regarded as a positive sign of a normal cardiovascular system, for it may occur in any form of heart disease.

SINUS TACHYCARDIA

The heart rate varies markedly in different mammals. In the elephant for example it is about 30 beats per minute in the rat it is close on 600. It is considerably slower in the hare than in the rabbit. On the whole the speed is inversely proportional both to the size and to the athletic endurance of the animal. In man the average heart rate is 72 beats per minute but there are wide limits of normality ranging between 40 and 100. The pulse is faster in children averaging 120 to 130 at birth and slowing gradually during childhood to reach about 80 at puberty. The more athletic the individual the slower the pulse as a rule and in well trained athletes resting figures of 45 to 50 are common. It follows that tachycardia may mean a heart rate faster than average faster than the upper limit of normality or faster than what is known to be normal for a particular individual.

Applied physiology. Tachycardia has always played an impressive part as a physical sign in general medicine. It has received weighty consideration in fevers in all forms of heart disease in shock and hæmorrhage in various chronic diseases such as pulmonary tuberculosis and indeed in almost every condition yet it can mean little unless its immediate cause is understood. This is not to decry tachycardia as a valuable sign but to emphasise that its significance depends upon its mechanism.

The speed of the sino auricular pace-maker is strongly influenced by the autonomic nervous system. Complete paralysis of the vagus may be produced within a minute by giving 2 to 3 mg. of atropine sulphate intravenously whereupon the heart accelerates to a speed of 130 to 160 per minute. The cardiac output per minute rises simultaneously but a fall in venous filling pressure which accompanies the tachycardia may counteract this effect (McMichael and Sharpey-Schafer 1944). The ventricular stroke volume is diminished even in those with higher outputs. Emotional tachycardia as in the anxiety states and also the tachycardia of convalescence appear to be due to diminished vagal tone.

Tachycardia may be due to a rise in pressure within the great veins and right auricle venous receptors initiating the Bainbridge reflex by which vagal tone is reduced. Under these circumstances the stroke volume may be maintained or increased the cardiac output per minute rising in proportion to the tachycardia or even higher. This mechanism operates during effort and in anæmia beri beri arteriovenous shunt anoxic pulmonary heart disease generalised active Paget's disease and pregnancy. The Bainbridge reflex is also responsible for the tachycardia so frequently seen in congestive failure.

The speed of the heart is also controlled by reflexes initiated by receptors in the aorta and carotid sinuses. When the blood pressure rises vagal tone is increased and the heart slows when it falls vagal tone is diminished and the heart quickens (Marey's law). This is the mechanism of tachycardia associated with conditions causing a transient rise of blood pressure.

such as acute nephritis and it is part of the mechanism controlling the tachycardia of low blood pressure states.

Anoxia may cause tachycardia by direct action on the central nuclei or possibly reflexly through the carotid sinus. Just what part it plays in the production of tachycardia in anæmia and cor pulmonale is uncertain. Thyroxin and fever have a direct stimulating action on the pace maker and so has adrenaline but the latter may also excite the carotid sinus slowing reflex by raising the blood pressure so that the heart rate may change but little. The elevated cardiac output which accompanies the tachycardia is also probably due in part to a direct action on the heart. In the case of adrenaline the cardiac output may rise when there is no change in heart rate or blood pressure (McMichael and Sharpey Schafer 1944).

Differential diagnosis From the clinical point of view sinus tachycardia must be distinguished from auricular flutter and from paroxysmal tachycardia. This is usually possible at the bedside. Sinus tachycardia varies in

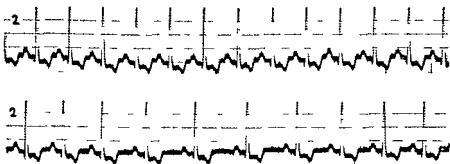


Fig. 403—Sinus tachycardia slowed by carotid sinus compression

rate from minute to minute or at least from hour to hour and it varies with emotion, effort and change of posture. Carotid sinus or eyeball compression and release result in gradual rather than abrupt slowing and quickening of the pulse respectively although changes may be difficult to detect with fast rates. In auricular flutter (and sometimes in paroxysmal auricular tachycardia) the rate is usually fixed neither varying spontaneously nor with emotion, effort or change of posture whilst on carotid sinus pressure slowing is abrupt often to half the rate 2:1 physiological auriculo-ventricular block being converted into a 4:1 relationship and on release reversion to the original rhythm is again abrupt and may not take place for several seconds. Even without so precise a clinical analysis the degree of slowing may yet be too gross for sinus tachycardia. In paroxysmal nodal and ventricular tachycardia the rate is also fixed and carotid sinus pressure either stops the attack abruptly as in 50 per cent of the nodal cases or has no effect whatever. If it is impossible to interpret the results of carotid sinus pressure clinically the problem may be solved by combining the manoeuvre with an electrocardiogram (fig. 403). It should be

explained that an electrocardiogram *per se* may not afford certain distinction between the e three rhythms although lead V_1 or CR1 greatly facilitates analysis

Effect on the heart Sinus tachycardia presents an important problem in relation to heart failure. Is it a causal factor or merely a reflection of cardiac embarrassment. Or is it part of a compensatory adjustment beneficial under the circumstances. Such questions are difficult to answer directly but the presentation of some of the relevant facts may help to clarify the issue. A normal heart tolerates any natural degree and duration of sinus tachycardia rates approaching 200 for example being common during violent exertion and persistent rates of 120 or so being endured for over 20 years in certain cases of Da Costa's syndrome without harmful results. On the other hand diseased hearts frequently develop congestive failure with heart rates of 130 to 200 in auricular flutter or paroxysmal tachycardia the effect being attributed to overwork and to fatigue resulting from insufficient diastolic rest. The tachycardia of the hyperkinetic forms of cardiovascular disorder (thyrotoxicosis anæmia anoxic pulmonary heart disease beri beri arterio venous aneurysm and generalised Paget's disease) is part of the physiological mechanism maintaining a high cardiac output and therefore performs a useful function but when the heart fails i.e. when it is overloaded the cardiac output falls and the tachycardia is wasted. Under such circumstances tachycardia reflects cardiac embarrassment and deprives the heart of diastolic rest. In the hypokinetic forms of heart failure such as those which may be seen in cases of hypertension and mitral stenosis tachycardia due to the operation of the Bainbridge reflex is a reflection of cardiac distress from the start and serves no useful purpose. In chronic constrictive pericarditis and to a lesser extent in high pressure pericardial effusion tachycardia may provide the only means of maintaining an adequate cardiac output for the stroke volume is strictly limited. In the active forms of carditis (rheumatic diphtheritic and Fiedler's) and in bacterial endocarditis the heart rate may be disturbed by local pathology fever toxæmia or (in diphtheria) by circulatory collapse and probably adversely affects the heart. On the whole it may be said that the heart tolerates sinus tachycardia which tends to deprive it of rest better than a high cardiac output and much better than a raised blood pressure both of which increase its work.

There is no treatment for sinus tachycardia itself but attention should be paid to its cause.

SINUS BRADYCARDIA

As already stated heart rates of 45 to 50 per minute are common in athletes. Some individuals irrespective of their physical training have a naturally slow pulse. Sinus bradycardia is a feature of certain diseases notably myxædema obstructive jaundice and aortic stenosis and it is not uncommon during convalescence from certain fevers especially influenza.

It also occurs when the blood pressure is raised rather suddenly as in acute nephritis the slowing being reflex through the sino aortic afferents and the vagus. It is a familiar sign of lesions that increase the intracranial pressure when it may be due to direct stimulation of central nuclei. Slowing of the pulse may be induced temporarily by carotid sinus or eyeball pressure as a transient event it occurs naturally in vaso vagal syncope.

The differential diagnosis between sinus bradycardia sino auricular block and heart block can usually be made at the bedside but electro cardiographic confirmation is advised. In sinus bradycardia the pulse quickens gradually with effort atropine or amyl nitrite in sino auricular block and sometimes in 2:1 heart block the rate doubles abruptly whilst in complete heart block the degree of acceleration is barely perceptible. Heart block may also be recognised by studying jugular pulsation (see page 120).

One of the consequences of sinus bradycardia is an increased ventricular stroke volume of sufficient degree to maintain a normal cardiac output per minute. When the heart rate is 40 the stroke volume approaches double the average normal the diastolic heart size is larger than usual (fig 4.04) and in time hypertrophy may occur. Such enlargement is physiological.

When the speed of the pace maker approaches 40 per minute it may become slower than the natural speed of impulse formation in the auriculo ventricular node in which event nodal rhythm occurs. As sinus arrhythmia is often associated with bradycardia it is more usual to see irregular examples of ventricular escape the A V node taking over whenever a pause is unusually long (fig 4.05). Nodal rhythm would supervene more frequently if the influences which retarded the sinus node did not also depress the A V node.

Sinus bradycardia is often associated with sinus arrhythmia sometimes with auricular ectopic beats and rarely with paroxysmal auricular fibrillation or flutter in elderly subjects. Vagal influences appear to be responsible

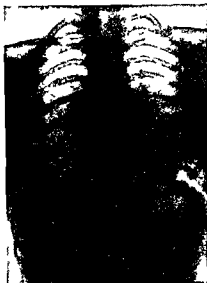


Fig 4.04—Relative cardiac enlargement due to sinus bradycardia

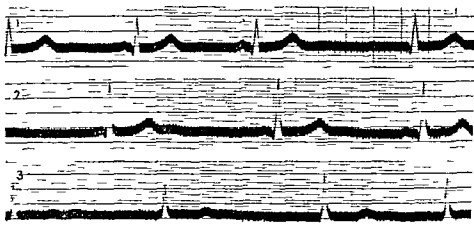


Fig 4 05—Nodal escape in sinus bradycardia

SINO AURICULAR BLOCK

There are three types of sino auricular block corresponding to similar varieties of A V block. First beats may be dropped irregularly the pauses being roughly equal to two normal intervals (fig 4 06) like the dropped beats of partial A V block with fixed prolonged P R interval. Second beats may be dropped more or less regularly the pauses being always less than two normal intervals like partial A V block with progressive lengthening of the P R interval until conduction fails—the Wenckebach type. Third there may be 2 : 1 sino auricular block every second beat being dropped this gives rise to a slow regular heart rate which doubles on effort or with atropine (fig 4 07). To understand these phenomena it is necessary to appreciate the fact that there is no electrocardiographic representation of the formation and discharge of the excitatory impulse at the sinus node, the first wave (P) of the electrocardiogram recording the passage of the impulse through the auricles so that failure of conduction between the S A node and the auricles can only be inferred.

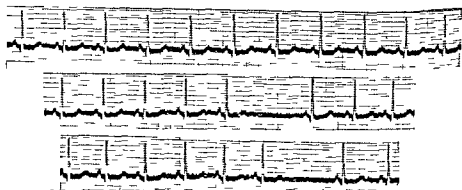


Fig 4 06—Sino auricular block showing irregular dropped beats



Fig. 407—No sinoauricular block: the rate doubles on effort

Sinoauricular block is usually encountered in normal individuals, the first two types being commonly associated with sinus bradycardia. It is a manifestation of increased vagal tone and may be abolished with atropine. When there is 2:1 block and a pulse rate of about 40 per minute, fluoroscopy may reveal cardiac enlargement due to the large stroke volume necessary to maintain a normal cardiac output. As with sinus bradycardia,

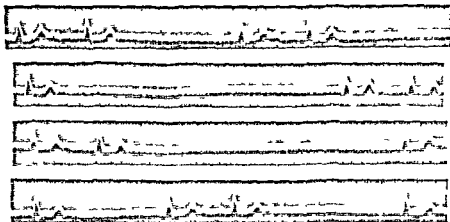


Fig. 408—Cardiac standstill occurring spontaneously in sinoauricular block
(By courtesy of Dr. R. J. M. and Dr. L. J.)

ventricular escape may occur and would probably be more common if the A-V node were not also depressed.

There are no symptoms of sinoauricular block *per se*, but occasionally short periods of cardiac standstill with dizziness or syncope may occur and appear to be due to bursts of extreme vagal activity (Fig. 408). They may be prevented by atropine. Attacks of this kind may be readily induced in susceptible individuals by carotid sinus pressure (Fig. 409).

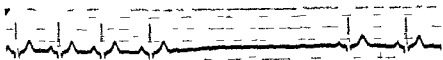
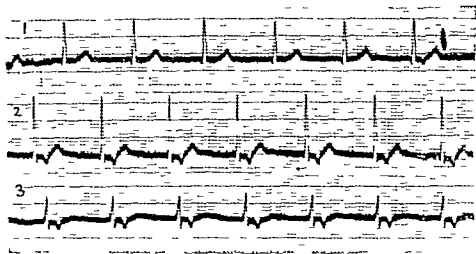


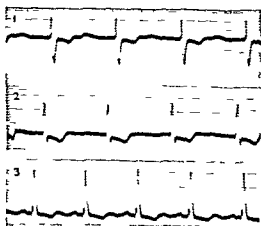
Fig. 409—Cardiac standstill due to carotid sinus compression



(a) An inverted P wave occurs after QRS



(b) Inverted An inverted P wave precedes QRS (Coronary sinus rhythm)



(c)—The P wave is invisible being buried in QRS (leads 1 and 2 lead 3 shows normal rhythm)

Fig 4 to—Nodal rhythm

NODAL RHYTHM

The sinus node is the pace maker of the heart only because its inherent rate of impulse formation and discharge is quicker than that of any other focus endowed with a similar capacity but if it is sufficiently depressed as by cooling some other focus may form its impulses at a faster rate and so become the temporary pace maker and in fact this function usually falls upon the auriculo ventricular node Under such circumstances auricular excitation is retrograde and the electrocardiogram usually shows an inverted (or deformed) P wave just after the QRS complex and a heart rate of 40 to 60 per minute (fig 4 10a) Sometimes however the P wave may precede (fig 4 10b) or coincide with the QRS complex or it may be absent altogether owing to retrograde block (fig 4 10c) Occasionally it may shift

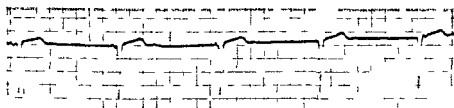


Fig 4 11—Shifting nodal rhythm

its position from moment to moment (shifting or sliding nodal rhythm fig 4 11) but if such graphs are examined critically some are seen to be examples of sinus bradycardia with frequent ventricular escape (so called wandering or shifting pace maker) and others reveal progressive lengthening of the period of retrograde conduction to the auricles until an auricular beat is dropped after which the P wave reverts to its initial position the cycle being repeated indefinitely (partial retrograde block)

Nodal rhythm may be discovered by chance in healthy individuals it may occur in active rheumatic diphtheritic and Fiedler's carditis it may be momentarily induced by carotid sinus pressure and it may follow thrombosis of the right coronary artery above the origin of the branch to the sinus node (this branch arises from the left coronary artery in 40 per cent of cases) but its only common cause is digitalis therapy

Nodal rhythm is under autonomic control the heart rate being slowed by vagal stimulation and accelerated by atropine and exercise (White 1915) It is a harmless rhythm change gives rise to no symptoms and requires no treatment When due to digitalis there is no need to stop the drug

HEART BLOCK

When any organic lesion or functional disturbance impedes conduction through the bundle of His or through both its main branches we may speak of heart block There are four grades prolonged P R interval

dropped beats partial block with fixed auriculo ventricular relationship and complete heart block

PROLONGED P R INTERVAL

As discussed on page 78 the upper limit of the normal P R interval should not exceed 0.22 second. In partial heart block it frequently measures 0.28 to 0.32 second. In extreme cases or when there is associated tachycardia electrocardiograms may show P coinciding with or even preceding the previous T wave (fig. 4.12).

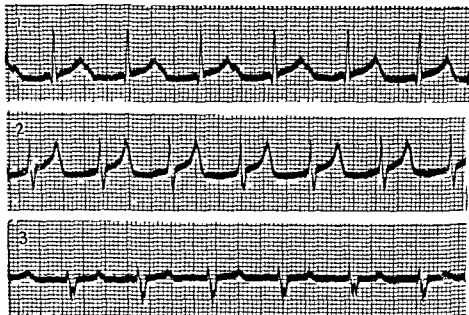


Fig. 4.12—Prolonged P R interval with P coinciding with the previous T wave

Prolongation of the P R interval may be transient or permanent or it may develop into a higher grade of block. As a transient phenomenon (when it may be abolished by the intravenous injection of 2 to 3 mg. of atropine sulphate) it is especially characteristic of any form of active carditis but it may also be due to digitalis to coronary thrombosis or to temporary nutritional changes from other causes and it may be induced by carotid sinus pressure. Permanent delay in conduction may result from an inflammatory scar involving the bundle of His as in old rheumatic heart disease or from ischaemic fibrosis.

Although partial heart block of this kind is essentially an electrocardiographic diagnosis it may be suspected clinically on occasions by noting delay between the auricular and ventricular components of cervical venous pulsation by observing a gap between a presystolic murmur and the first heart sound in cases with mitral stenosis or by detecting Cannon waves

in the neck (p. 122). Its practical importance lies in its value as a sign of active rheumatic carditis.

No special treatment is required.

PARTIAL HEART BLOCK WITH DROPPED BEATS

In a slightly higher grade of partial heart block, conduction through the bundle of His fails altogether from time to time so that ventricular beats are dropped. In the type first recognised by Wenckebach (1899) the P-R

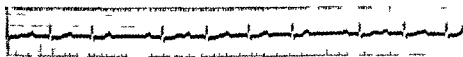


Fig. 4.13—Partial heart block with dropped beats (Wenckebach type)

interval shortens considerably after a beat is dropped but subsequently lengthens progressively from cycle to cycle until conduction again fails (fig. 4.13). In another type (Hay, 1906) the P-R interval is fixed and beats are dropped irregularly and unpredictably.

The condition may be suspected clinically, but cannot be so distinguished from partial sino-auricular block, nor from pauses following extremely premature and therefore inaudible ectopic beats. It is commonly transient and recovers spontaneously, but occasionally progresses to complete heart block.

PARTIAL HEART BLOCK WITH FIXED 4:1 RELATIONSHIP

Relatively stable forms of partial heart block may be encountered, usually with a 2:1 auriculo-ventricular relationship (fig. 4.14), but occasionally

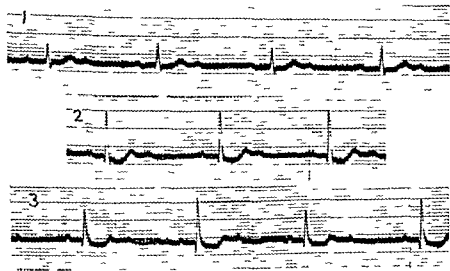


Fig. 4.14—2:1 heart block

with 3 : 1 or even 4 : 1 A : V ratios. These usually progress to complete heart block; they are much less common in active carditis than in ischaemic cases.

Clinically 2 : 1 heart block has to be distinguished from sino auricular block from sinus bradycardia with a heart rate of about 40 per minute from nodal rhythm and from complete A : V dissociation. Failure to quicken appreciably with effort or atropine excludes sino auricular block and sinus bradycardia (and usually nodal rhythm). If isolated auricular waves can be detected in the veins of the neck their regular timing may distinguish 2 : 1 block from complete A : V dissociation.

COMPLETE HEART BLOCK

Etiology. Complete auriculo ventricular dissociation is very rare in active rheumatic carditis but less so in diphtheritic carditis; it may be induced by digitalis especially in cases of auricular fibrillation and has been caused by hæmorrhage into the bundle of His from trauma or asphyxia and by primary or secondary neoplasm. About 10 per cent of cases are congenital and may be associated with ventricular septal defect. As a rule however complete heart block is associated with ischaemic or hypertensive heart disease with syphilitic aortitis or with extensive calcification of the aortic cusps or mitral ring in elderly atherosclerotic subjects and is due to a fibrotic or calcified lesion in the bundle of His, or in both its main branches.

Clinical features. A : V dissociation is four times more common in males than in females and 84 per cent of cases occur in patients over 50 years of age (Campbell 1944). It is usually permanent but under special circumstances may be transient or even paroxysmal (Lawrence and Forbes 1944). It is characterised by an extremely slow heart rate by a water hammer or collapsing pulse by elevation of the venous pressure by cervical venous pulsation unrelated to ventricular contraction by audible independent auricular sounds by the occurrence of Cannon waves in the neck and varying intensity of the first heart sound by general enlargement of the heart and by syncopal attacks of a special kind. It is proved electrocardiographically (figs 4 15a and b).

Whilst the pulse rate is usually about 28 to 36 per minute based on the inherent rate of impulse formation of the idio ventricular pace maker distal to the block in the bundle of His it may be so slow as to induce a state of continual faintness (fig 4 15b) as in the case originally described by Spens (1793) in which it fell to 9 beats per minute. At the other extreme complete A : V dissociation may be seen with a ventricular rate of over 100 the ventricles sometimes beating more rapidly than the auricles (fig 4 16). On the whole rates are faster when QRS is normal in width slower when the QRS resembles left or right bundle branch block (Kay 1948). Idio ventricular pace makers are little affected by stimuli which influence the S : A and A : V nodes so that the pulse rate usually remains remarkably

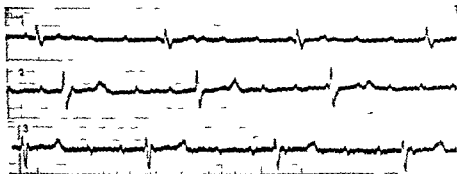


Fig 4 15 (a)—Complete heart block Ventricular rate 18 beats per minute

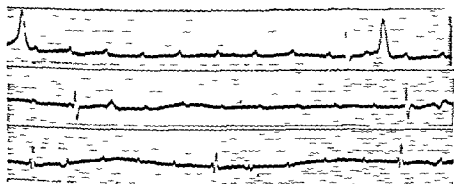


Fig 4 15 (b)—Complete heart block Ventricular rate 10 beats per minute

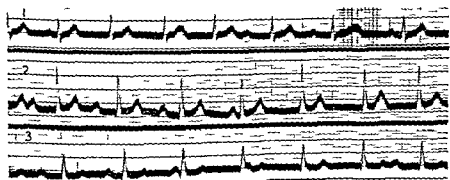


Fig 4 16—Complete A-V dissociation with the ventricles beating faster than the auricles

constant in complete heart block. In transient or paroxysmal cases however in which a functional element may be present, temporary restoration of sinus rhythm may accompany fever as in the case described by Gerbezius in 1719 (Major 1932).

A high systolic blood pressure is usual, and is due to the large ventricular stroke volume. Owing to associated vasodilatation however the pressure is not well maintained but tends to fall away rapidly in diastole giving rise to a collapsing pulse and to a rather low diastolic blood pressure.

Under favourable circumstances inspection of cervical venous pulsation may reveal auricular waves (a waves) independent of ventricular events (c and v waves), as noted by Stokes (1846). Simultaneously may be heard the faint sounds of isolated auricular contractions (the semi beats of Stokes) either at the apex beat or down the left border of the sternum.

Venous cannon waves occur when the P wave falls between QRS and T i.e. when the auricle contracts against a closed tricuspid valve and are easily recognized by their abrupt quality, high amplitude and variability. Changing intensity of the first heart sound is equally characteristic: the loudest sounds are heard when the P-R interval is around 0.10 to 0.12 second, auricular contraction then forcing the mitral cusps wide open just before ventricular systole (Levine 1948).

Cardiac enlargement is usually more conspicuous than that seen in sino-auricular block or in sinus bradycardia but is of the same quality unless the size and shape of the heart are altered by other effects of the underlying disease process.

The cardiac output can only be maintained by a large stroke volume propelled with great force. Diastolic distension is favoured by a compensatory rise in venous pressure and this must be very considerable during effort. The early development of congestive failure is readily understood.

Stokes Adams attacks—Syncope due to ventricular asystole (Stokes Adams attacks) occurs in about 50 per cent. of cases and is especially common when partial block becomes complete. Loss of consciousness is abrupt without warning. If standing the patient collapses and lies limp, still pale and pulseless with fixed dilated pupils—as if dead, breathing however continues. If the attack lasts long enough i.e. for more than 10 seconds or so twitchings commence and may progress to convulsions and if ventricular asystole continues for more than 2 or 3 minutes recovery is rare. As a rule however ventricular beating is resumed after a few seconds, consciousness returns abruptly and a vivid flush ensues. When an attack occurs in bed the lack of warning, short duration of unconsciousness and abrupt return of full possession of the faculties may prevent a dull patient from being aware of the fit and he may only notice the flush. The sequence of events both symptomatically and objectively is so characteristic as to make the diagnosis probable on the history alone—a point of some importance in patients with paroxysmal block who may present themselves with

normal sinus rhythm. In such cases carotid sinus pressure may provoke an attack or induce paroxysmal heart block (fig. 4 17).

Physiologically Stokes Adams attacks are due to depression of a potential or established idio ventricular pace maker in cases of complete heart block the ventricles stand still while the auricles continue to beat. They are apt to occur when partial block becomes complete either because such an event is usually associated with some depressive influence on conduction which may also depress ventricular pace makers (even though considered beyond vagal control) or because idio ventricular pace makers are by nature initially sluggish. When complete block is well established attacks

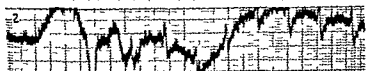
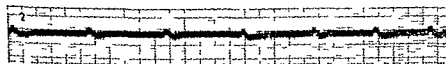


FIG. 4 17—Stokes Adams fit artificially provoked in a patient with paroxysmal complete heart block.

may still occur but are less common. The abrupt loss of consciousness depends upon sudden total failure of cardiac output. Twitching is due to cerebral anoxia and is not seen in short attacks. Convulsions are of two types, one being an exaggeration of anoxic twitching, the other occurring after restoration of ventricular action and synchronising with the flush (Formijne 1938). In the second type convulsions and flushing appear to be due to carbon dioxide depletion in the blood stagnant in the lungs during the phase of asystole with continued respiration and to vasodilatation resulting from accumulation of tissue metabolites so that when ventricular beating is resumed blood rich in oxygen but containing practically no carbon dioxide is thrown abruptly into a widely dilated vascular bed. More often a period of apnoea follows the attack with or without subsequent Cheyne Stokes breathing (Criffith 1921). Apnoea of course may also occur towards the end of long periods of ventricular asystole when it is due to failure of the respiratory centre resulting from profound cerebral anoxia.

An important complication of Stokes Adams attacks is paroxysmal ventricular tachycardia or fibrillation (Parkinson Papp and Evans 1941). In such cases it may be impossible to determine clinically whether uncon-

sciousness is due to asystole or to ventricular fibrillation. It is probable that many deaths are due to the supervention of such rhythm changes rather than to asystole.

Prognosis Congenital and transient cases do relatively well unless the disease responsible is serious for other reasons. The outlook in paroxysmal and acquired permanent cases however is poor, life expectancy averaging $4\frac{1}{2}$ years (Graybiel and White 1936; Campbell 1944). Those with a history of Stokes-Adams fits have a much worse prognosis than those without, the majority of them dying suddenly. Those without fits usually die from congestive heart failure.

Treatment The most effective prophylactic treatment for faintness or syncope is the oral administration of ephedrine $\frac{1}{2}$ grain (32 mg) t d s. If attacks are frequent and the patient bedridden, adrenalin 0.5 mg (8 minims or 0.5 ml of a 1:1000 solution) should be injected subcutaneously and repeated every two to six hours. Sublingual nor-adrenalin is also helpful. Both ephedrine and adrenalin prevent undue depression of the ventricular pace-maker and encourage the heart to beat a trifle faster. It is sometimes said that idio-ventricular rhythm cannot be influenced by any of the drugs or manœuvres that are known to effect the sinus node. This is not always strictly true, but changes are admittedly slight. Effort for example may quicken the ventricular rate in complete heart block, the adrenergic drugs, fever, and even atropine may also do so. In treatment, however, atropine is valueless alone, although it may enhance the effect of adrenaline. Barium chloride had a vogue, its action depending upon its power to excite ventricular ectopic beats and so to prevent ventricular standstill, but this is a poor substitute for the physiological benefit provided by ephedrine. In paroxysmal cases, when some functional disturbance must be postulated, inhalations of amyl nitrite may abort attacks (Lawrence and Forbes 1944).

A problem arises when repeated seizures are partly due to paroxysmal ventricular tachycardia or fibrillation, for if it is uncertain whether unconsciousness is due to asystole or to fibrillation, the administration of adrenaline becomes hazardous, as the drug encourages the latter rhythm change. For this reason electrocardiographic analysis is advised whenever possible.

Treatment of the primary cardiac condition may help. This applies especially to the rare transient cases associated with active carditis or myocardial infarction, and to permanent cases associated with syphilitic aortitis. Very rarely a small gumma may interrupt the conducting pathway, and the resulting block may be cured with iodides (Major 1923).

If congestive heart failure calls for digitalis therapy, the drug should not be withheld on account of coincident heart block, but should be administered with caution. Massive and intravenous doses should be avoided, but digitalis leaf 3 grains (0.2 G) t d s on the first day, 2 grains (0.13 G) t d s on the second, and 1 grain (0.065 mg) t d s thereafter until an adequate effect is obtained or until signs of intoxication occur, is usually safe. Should a Stokes-Adams fit appear to be provoked, the drug must be discontinued.

BUNDLE BRANCH BLOCK

Although bundle branch block is not strictly a disorder of rhythm it may be discussed here conveniently on account of its close pathological relationship to other forms of conduction defect

Anatomy Bundle branch block occurs when some organic lesion interferes with conduction through one or other of the two main branches of the bundle of His. As may be seen from figure 401 the main bundle after piercing the membranous septum divides into two one branch passing down each side of the muscular interventricular septum just beneath the endocardium and spreading out fan wise distally the left branch may subdivide into anterior and posterior divisions in the lower half of the septum (Mahaim 1931). The A V node bundle of His and posterior division of the left bundle branch receive their blood supply from perforating septal arteries arising from the posterior descending branch of the right coronary artery the right bundle branch and the anterior division of the left are supplied by perforating septal branches of the left anterior descending coronary artery (Gross 1921). Considerable variations occur however especially as vital reactions to ischaemia

Nomenclature When the left bundle branch is interrupted the excitatory process reaches the right ventricle first through the relatively normal right bundle branch and spreads throughout that chamber before passing across to the left. The right ventricle therefore contracts first. The electrocardiogram described and illustrated fully in Chapter III shows a wide QRS complex measuring from 0.11 to 0.18 second the main deflection of which is usually upright in lead 1 and downward in lead 3 with marked slurring or notching and followed by a conspicuous T wave usually in the opposite direction. Right bundle branch block (Wilson *et al* 1934) is characterised by widening of the initial ventricular deflection to 0.11 to 0.14 second by late slurring of QRS — usually best seen in S_1 — and by an upright T wave in lead 1 (page 93). That the first type of graph described represents left bundle branch block has been proved by the reconstructed vectorcardiograms (monocardiograms) of Mann (1931) by the electrocardiographic discoveries of Wilson and his colleagues (1932) by kymographic and polygraphic studies revealing delayed left ventricular events (Wolferth and Margolies 1935) by experiments on revived human hearts in normal position in which one or other bundle branch has been cut (Kountz 1936) and by simultaneous electrocardiographic phonocardiographic and polygraphic records demonstrating and analysing ventricular asynchronism (Braun Menendez and Solari 1939). The detailed histological work of Mahaim (1931) which at first appeared to support the original view in which the nomenclature for left and right bundle branch block was reversed has been ably reviewed by Yater (1938) who presented extensive histopathological evidence of his own and concluded that the bilateral lesions invariably demonstrable rendered reliable interpretation difficult

but that on the whole the findings supported the new terminology. Finally, the clinical facts cannot be disregarded: left bundle branch block is commonly seen in lesions involving the left side of the heart whereas right bundle branch block is usually associated with enlargement of the right ventricle. This general principle was recognised by Tung and Cheer (1933) and by Bayley (1934).

Etiology Left bundle branch block is usually due to hypertensive heart disease, ischaemic heart disease or aortic valve disease; right bundle branch block to mitral stenosis, atrial septal defect or massive pulmonary embolism. Either form may occur in active rheumatic, diphtheritic or other

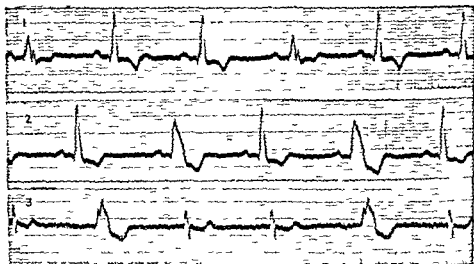


Fig. 4 18—Alternating left bundle branch block

form of carditis in any disease affecting the heart as a whole such as thyrotoxicosis and fibrosis of the myocardium of known or unknown etiology and as a result of any local lesion such as neoplasm. Partial forms are common and tend to progress; on the other hand both left and right bundle branch block may be transient, paroxysmal or even alternating (fig. 4 18) sometimes in association with paroxysmal tachycardia, auricular flutter or fibrillation, sometimes during an episode such as acute myocardial infarction, congestive heart failure or massive pulmonary embolism but also spontaneously. Right bundle branch block is sometimes found in otherwise healthy individuals even in youth.

Clinical features Clinically left bundle branch block may be suggested by presystolic gallop rhythm in the absence of ventricular distress and right branch block by wide splitting of the second heart sound. In many cases however no such clue is afforded and its existence is only discovered electrocardiographically. When the heart is enlarged and it is uncertain which chamber is mainly involved the presence of left or right bundle

branch block points strongly to the homolateral ventricle. Left bundle branch block provides convincing proof of serious heart disease but right bundle branch block must be interpreted more cautiously. Neither form is influenced by digitalis, atropine or by any of the adrenergic or cholinergic drugs.

Prognosis. The average life expectancy for cases of bundle branch block in general has been estimated at 3 years (Campbell 1944) but it should be clearly understood that in any given patient the prognosis is that of the underlying heart disease and is not influenced by the conduction defect. Again if right bundle branch block is found in an otherwise normal individual the outlook does not differ from normal controls (Wood, Jeffers and Wolferth 1935).

ECTOPIC BEATS

Ectopic beats are premature systoles induced by the discharge of some ectopic impulse forming focus situated anywhere in auricular, nodal or

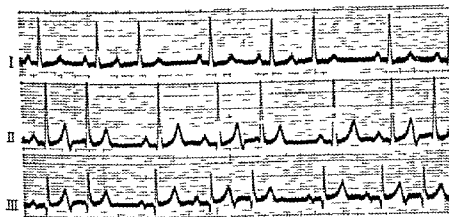


FIG. 4-19—Auricular ectopic beats

ventricular tissue. They are necessarily premature because all potential impulse forming foci are otherwise discharged by the excitation which reaches them from the sinus node.

Physiology. In the auricular type (fig. 4-19) the P wave is abnormal in shape or direction according to the site of the ectopic focus and to the direction in which the impulse flows over the auricles. In these cases the partially charged sinus node is discharged when the impulse reaches it so that the compensatory pause following the ectopic beat is slight being equal to a normal cycle plus the interval between the onset of the ectopic and the arrival of the retrograde excitatory process at the S-A node. The timing of the heart beat is permanently altered. The ventricular complex is usually normal but may be slightly deformed as a result of a functional defect in

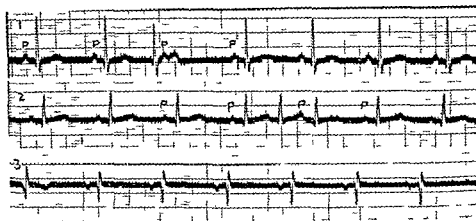


Fig 4 20—Nodal ectopic beats. Slight deformity of QRS is due to fatigue block. In lead 1 the P wave immediately after the ectopic is blocked. In lead 2 the nodal ectopic is interpolated. In both there is retrograde block.

conduction If an auricular ectopic beat is very premature it may be blocked altogether.

Nodal ectopic beats (fig 4 20) are premature beats arising in any part of the auriculo ventricular junctional tissue. The QRS complex is normal or slightly deformed as described above, but the P wave is inverted and occurs just before, during, or just after the QRS complex, according to the more proximal or more distal site of the ectopic focus, and to the degree of resistance opposed to retrograde conduction. Discharge of the sinus node (unless there is retrograde block) again prevents a full compensatory pause.

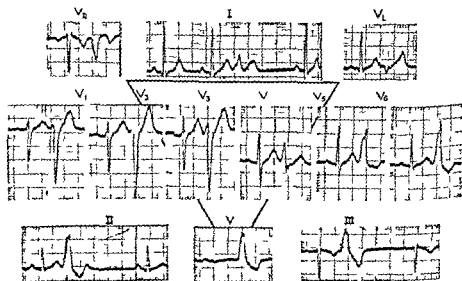


Fig 4 21—Right ventricular ectopic beats.

Ventricular ectopic beats are characterised by a full compensatory pause for the sinus node is not discharged by the premature impulse owing to retrograde block (physiological) in the bundle of His or to natural delay in retrograde conduction and so continues to function at its usual time. Its first discharge after the ectopic however is blocked by the refractory state of the ventricles and so there is a pause until its second discharge. The final timing of the heart beat therefore remains unchanged. Electrocardiographically a ventricular ectopic beat resembles a bundle branch block

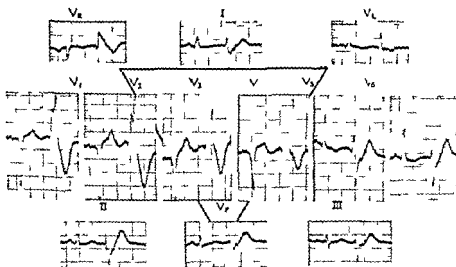


Fig. 4.22—Left ventricular ectopic beats causing coupling

complex QRS being widened and notched and T being prominent and usually in the opposite direction. When the deflection is like left bundle branch block the ectopic focus lies in the right ventricle (fig. 4.21) when QRS is like right bundle branch block the ectopic focus lies in the left ventricle (fig. 4.22). There are many variations however depending upon the exact site of the irritable focus (Barker *et al.* 1930; Kountz 1936).

Premature beats have a smaller stroke volume than normal and if very premature may not be perceptible at the wrist or audible with a stethoscope. The beat which follows is fuller than usual and is appreciated by the patient as a hard thump. This is a matter of cardiac filling: the earlier the ectopic beat the emptier the heart, the longer the compensatory pause the fuller the heart. The blood pressure varies directly with the amplitude of the pulse.

Clinical diagnosis. Clinically, ectopic beats must be distinguished from other irregularities especially from auricular fibrillation and from partial heart block with dropped beats. Whilst this may be easy in the majority of cases confusion arises with multiple auricular ectopic beats which may be indistinguishable from auricular fibrillation and with inaudible imper-

ceptible or blocked ectopic beats which mimic partial heart block with dropped beats. Alternate ectopic beats or coupled beats may be confused with S A block when very premature with a dicrotic or bisferiens pulse, or even with pulsus alternans. If there is any doubt the effect of effort, amyl nitrite or of 1 mg. of atropine sulphate should be determined. Ectopic beats usually disappear as the heart quickens and may be exaggerated as it slows down again.

Etiology. Experimentally ectopic beats may be produced by electrical stimulation of any part of the heart. Certain drugs notably digitalis, barium chloride and adrenaline may produce them. Excessive use of tobacco occasionally seems responsible. They are common in pregnancy. Whilst almost any state of ill health may be blamed for their occurrence, no common factor has been discovered and in the majority of cases there is no evidence of structural disease of the cardiovascular or other systems. Occasionally, however, auricular ectopic beats may herald auricular fibrillation especially in mitral stenosis and thyrotoxicosis. Under certain circumstances also, ectopic beats are probably due to organic disease for example their occurrence during the course of diphtheria may be due to toxic carditis but as innocent ectopic beats are common enough after simple streptococcal tonsillitis and indeed during convalescence from any fever it is impossible to draw any conclusion from their presence. Again ectopic beats following coronary thrombosis are probably significant, and to be explained by irritable foci set up by ischaemia but as they are equally common in conditions which may simulate myocardial infarction e.g. gall bladder disease, diaphragmatic hernia, upper abdominal catastrophes, acute anxiety states and the like they are of no diagnostic value. On the whole therefore it is wise to assume the innocence of ectopic beats under any conditions and to judge organic disease on other grounds.

Treatment. Many patients are unaware of premature systoles, others may seek relief from palpitations. Treatment includes fresh air, exercise and a healthy physiological life. Of drugs potassium bromide 10 grains (0.65 G) t.d.s., phenobarbitone $\frac{1}{2}$ grain (32 mg) t.d.s. or quinidine 5 grains (0.32 G) t.d.s. may prove effective. Alternate ectopic beats (coupling) due to digitalis provide good grounds for stopping the drug or reducing its dose. Potassium salts are efficient (Dampson and Anderson 1932, Castleden 1941) but the large dose usually required is not without danger of sudden death and may provoke symptoms as unpleasant as the palpitations, chiefly nausea and vomiting, the chloride or acetate is employed as a 10 to 20 per cent aqueous solution and may be given by mouth in doses of 2 to 6 G three or four times a day. Larger doses are not advised and when the maximum recommended is being administered the patient should be confined to bed especially if there is underlying organic heart disease. Reassurance is important, and should be unconditional and convincing for it should be remembered that ectopic beats rarely constitute a complaint except in those prone to morbid anxiety.

EXTRASYSTOLES

The term extrasystole should be reserved for interpolated ectopic beats (fig 4 23) These are true extra heart beats and occur when the impulse from the sinus node immediately after the ectopic beat manages to excite the ventricles They are usually ventricular but may be nodal (fig 4 20)

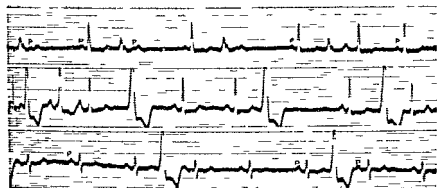


Fig 4 23—Interpolated ventricular ectopic beats

PAROXYSMAL TACHYCARDIA

When ectopic beats occur in rapid and regular succession from the same focus one may speak of paroxysmal tachycardia The name was introduced by Bouveret in 1889 The ectopic focus may be supraventricular (auricular or nodal) or ventricular The electrocardiographic complexes in the three types are precisely the same as those in the three types of ectopic beat

The patient usually complains of attacks of palpitations characterised by the abruptness of their beginning and end by the rapidity and regularity of the beats and by the relative well being of the patient (Cotton 1867) Until an attack is witnessed the diagnosis rests upon an accurate history Experience shows that most careful cross examination is required to establish the true sequence of events It is not enough to determine that the onset is sudden it is necessary to be sure it is abrupt that the full velocity of the attack is reached immediately in the space of one beat that from no sensation whatever maximum palpitation develops within one second To assess the rate and rhythm it is helpful to ask the patient to represent them by tapping with his finger The manner in which the attack ends may be more difficult to establish some patients become accustomed to the palpitations and gradually fail to perceive them others pass from a true paroxysm to sinus tachycardia without appreciating the change and their description of the end refers to the gradual slowing down of the sinus rhythm

Attacks may last from a few seconds to several weeks but are usually

measured in hours and rarely exceed three days. The speed ranges between 110 and 250 per minute, but is between 140 and 240 in 90 per cent of cases, and between 150 and 200 in 50 per cent (Campbell 1947). Occasionally however much faster rates have been recorded. For instance in one of Bouveret's cases the heart rate was 300 per minute. If the heart is normal as it is in 62 per cent of the supraventricular variety there are usually no other symptoms apart from those provoked by anxiety, but if the attack is unduly prolonged or the heart rate exceptionally rapid congestive failure or angina pectoris may occur. If the heart is abnormal however as it is in 80 per cent of the ventricular variety the rapid development of congestive heart failure is common. With very rapid rates syncope is said to occur and in ischaemic heart disease, status anginosus. Physiologically the effects depend upon the functional capacity of the heart to increase its output with tachycardia and on its ability to stand up to the extra work imposed with minimal rest. At any given moment there must be a critical rate above which the cardiac output falls.

SUPRAVENTRICULAR PAROXYSMS

As just indicated both paroxysmal auricular and nodal tachycardia are most commonly encountered in healthy individuals and have little more significance than ectopic beats or spontaneous fluttering of somatic muscle. They are fifteen times more common than ventricular paroxysms. When attacks occur in patients with heart disease the prognosis is not so good and depends upon the nature and severity of the cardiac lesion and the

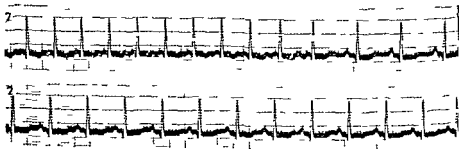


Fig. 4.24—Paroxysmal auricular tachycardia terminated by means of mechohn

speed and duration of the paroxysm. Even so the mortality rate is only about 1 per cent.

A clinical diagnosis may be accepted if the spontaneous or induced beginning or end of an attack is proved to be abrupt; if the heart rate during a paroxysm exceeds 150 per minute and does not vary with effort, change of posture, atropine, amyl nitrite, carotid sinus (or eyeball) pressure, prostigmine or mechohn; if any such measure terminates the paroxysm; if the duration of attacks is a matter of hours rather than one of minutes, days or

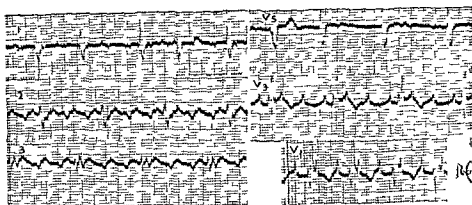
weeks if the patient is relatively young i.e. under 40 years of age or was so when he had his first attack if paroxysms have continued with variable frequency for more than five years and if there is no evidence of organic heart disease or thyrotoxicosis. Electrocardiographic proof however which may require a record of the beginning or end of an attack should be obtained whenever possible. Although only a rare chance will enable the onset to be registered the end may be recorded in over half the cases by means of a continuous tracing while the attack is terminated by carotid sinus pressure or mecholin (fig. 4 24). If the attack is not terminated such



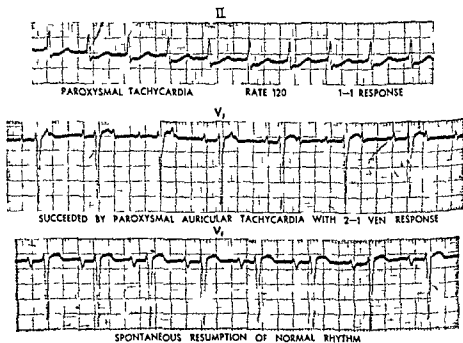
Fig. 4 25—Paroxysmal atrial tachycardia blocked by carotid sinus compression

measures may yet serve to differentiate paroxysmal tachycardia from sinus tachycardia and from auricular flutter for in paroxysmal tachycardia the heart rate is rarely altered whereas in sinus tachycardia it is slowed and in auricular flutter it is often abruptly halved. Occasionally however carotid sinus pressure may block paroxysmal atrial tachycardia (fig. 4 25).

Evans (1944) first presented evidence based on lead CR₁ suggesting that many cases which would ordinarily be interpreted as 2 : 1 auricular flutter might really be examples of paroxysmal atrial tachycardia with 2 : 1 A : V block and that these two conditions were essentially the same. Certainly paroxysmal atrial tachycardia may show varying degrees of A : V block (figs. 4 25 and 4 26a and b) and the auricular waves may be slowed by means of quinidine (figs. 4 27a and b) in the same way as flutter. There is also no doubt that the same patient may show all varieties of auricular rhythm suggesting that they all depend upon a similar mechanism and that the occurrence of auricular ectopics before or after a major attack (fig. 4 28) offers an obvious clue as to their essential nature. Indeed Prinzmetal (1950) has now provided convincing evidence not only of the unity of paroxysmal atrial tachycardia and flutter but also of auricular ectopic beats and auricular fibrillation all four disturbances of rhythm depending upon the presence and behaviour of an ectopic irritable focus. Nevertheless the clinical differences between paroxysmal tachycardia and flutter (not to mention auricular fibrillation and ectopic beats) are con-

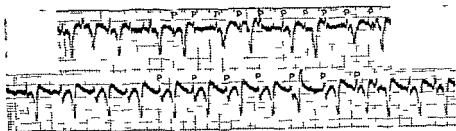


(a)

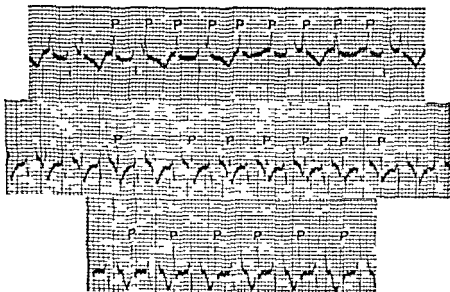


(b)

Fig 4 26 (a) (b)—Two cases of paroxysmal auricular tachycardia showing varying degrees of spontaneous A V block



(a)



(b)

Fig 4 27 (a) (b)—Two cases of paroxysmal auricular tachycardia slowed by means of quinidine

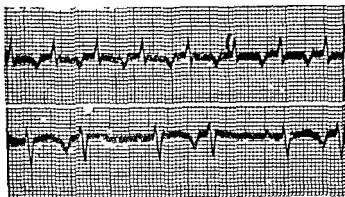


Fig 4 28—Paroxysmal auricular tachycardia followed by auricular ectopic beats

siderable (Campbell 1945) and their separate identities should be preserved. Similarly there can be no thought of not maintaining the separate identities of ventricular ectopics, ventricular tachycardia, and ventricular fibrillation, yet all three must depend on a similar physiological mechanism.

PAROXYSMAL NODAL TACHYCARDIA

Nodal paroxysms may be difficult to distinguish from auricular tachycardia when the rate is fast, unless the beginning of an attack can be recorded (fig. 4 29a). At slower rates the electrocardiogram resembles fast



Fig. 4 29 (a)—Paroxysmal nodal tachycardia beginning with a nodal ectopic beat

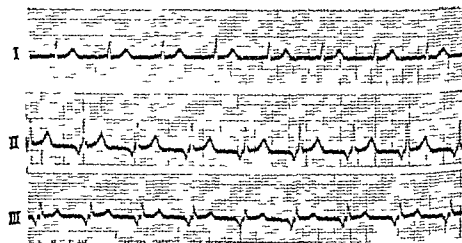


Fig. 4 29 (b) Abnormal nodal type of rhythm with a rate of 90 beats per minute. The pacemaker is either in the A V node or in adjacent auricular tissue such as the mouth of the coronary sinus.

nodal rhythm. Should an inverted P wave precede QRS, however, it may be impossible to determine whether the ectopic focus is in the A V node or in adjacent auricular tissue as in the slower form known as coronary sinus rhythm (fig. 4 29b).

Nodal tachycardia is commonly innocent and responds particularly well to carotid sinus pressure and to cholinergic drugs

INTRICULAR PAROXYSMS

Paroxysmal ventricular tachycardia is relatively rare is usually associated with organic heart disease in patients between the ages of 40 and 70 and is

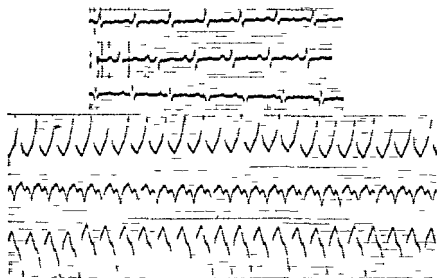


Fig 4 30—Paroxysmal ventricular tachycardia

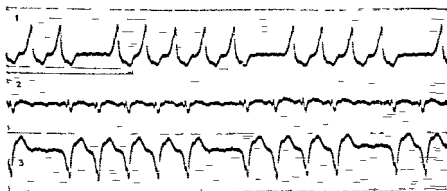


Fig 4 31—Auricular flutter with left bundle branch block

twice as common in men as in women. It tends to arise in a badly damaged heart as in heart failure from hypertension or from aortic valve disease; it may follow myocardial infarction or succeed a Stokes-Adams fit; occasionally it is due to digitalis; in about 20 per cent of cases it is innocent.

It has the same clinical features as the supraventricular variety apart from the circumstances in which it occurs and its lack of response to carotid sinus pressure and to the cholinergic drugs moreover it is more frequently followed by congestive heart failure and sometimes by ventricular fibrillation and sudden death. The prognosis is correspondingly grave.

Proof of the nature of the attack is obtained by electrocardiography (fig 4 30) but difficulty may arise when supraventricular paroxysms or auricular flutter are complicated by previously established or functional bundle branch block (fig 4 31).

TREATMENT

Supraventricular paroxysms may be terminated by some mechanical trick already known to the patient such as holding the breath, by carotid sinus or eyeball pressure in about 50 per cent of cases and by the cholinergic drugs in 75 per cent. Devices discovered by the patient include the adoption of some particular posture drinking cold water forced breathing or breath holding compression of the abdomen and self induced vomiting.

The carotid sinus is located at the bifurcation of the common carotid artery at the level of the superior border of the thyroid cartilage. It should be firmly compressed for several seconds against the bodies of the cervical vertebræ by means of the observer's thumb first on one side then on the other but never together. Bilateral eyeball compression is also carried out with the thumbs should be sufficiently forceful to cause pain and should be maintained for 3 to 5 seconds.

Of the cholinergic drugs mecholin (acetyl beta methylcholine) is the most successful and prostigmine the least unpleasant in the doses employed. doryl (carbo amino acetylcholine) is less effective, and acetylcholine itself too drastic besides being technically difficult owing to its rapid destruction in the bloodstream. Mecholin should be given intramuscularly or subcutaneously in a dose of 10 to 20 mg and may be expected to work in about five minutes. prostigmine may be administered intravenously or intramuscularly in a dose of 1 to 2 mg and has its maximum effect in about half an hour. Side effects include urgent micturition and defæcation colic vomiting sweating flushing and faintness but these are absent or slight with 1 to 1.5 mg of prostigmine and rarely severe with 10 mg of mecholin. Should they prove too unpleasant and the object of the drug has not been achieved they may be abolished at once by injecting 1 to 2 mg of atropine sulphate intravenously but this is obviously not advised unless absolutely necessary.

Should the cholinergic drugs fail to restore normal rhythm quinidine or even digitalis may be successful. In resistant cases of auricular tachycardia digitalis may be used to maintain 2:1 or a greater degree of A-V block in order to protect the ventricles. Bromide and phenobarbitone help to allay anxiety and congestive heart failure if it occurs should be treated by the usual methods.

Ventricular paroxysms do not respond to the above measures but may often be terminated by injecting 20 ml of a 20 per cent solution of magnesium sulphate intravenously (Szekely 1946) or by quinidine 5 grains (0.32 G) intravenously or in oral doses of 5 grains (0.32 G) two hourly to a maximum of 40 grains (2.6 G) in one day followed by 10 grains (0.65 G) two hourly for four doses on the second day if the first attempt fails and then by 15 grains (1 G) two hourly for three doses on the third day if necessary. Intravenous procaine in doses of 5 to 10 ml of a 1 to 2 per cent solution may also be tried.

To prevent both types of attack quinidine 3 to 5 grains (0.2 to 0.32 G) t d s may be continued indefinitely if well tolerated.

PAROXYSMAL TACHYCARDIA ASSOCIATED WITH PRE EXCITATION

The condition first described as physiological bundle branch block with short P R interval (Wolff Parkinson and White 1930) is probably due to premature excitation of one or other ventricle usually the right resulting from an anomalous connexion between the A V node or right auricle and the right ventricle (Holzman and Scherf 1932). Electrocardiography shows



Fig 4.32 (a)—Pre excitation

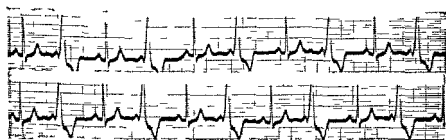
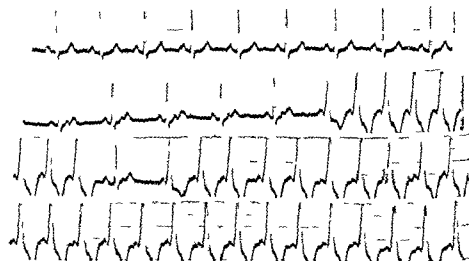


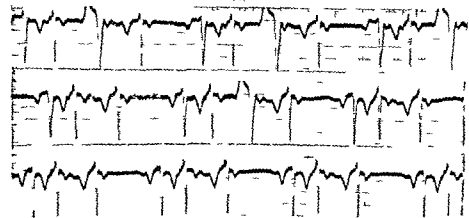
Fig 4.32 (b)—Alternating pre excitation



(a)



(b)



(c)

Fig 4 33—Wolff Parkinson White Syndrome showing
 (a) Paroxysmal tachycardia with normal QRS complexes (b) Paroxysmal tachycardia
 with widened QRS complexes (c) Auricular re entry following the major attack

widening of the QRS complex as in bundle branch block but at the expense of the P R interval which is shortened proportionally the P S interval as measured from the beginning of P to the end of the QRS complex remaining unchanged (fig 4 32a) The appearances usually resemble left rather than right bundle branch block The anomalous pathway may be through the bundle of Kent (Wood Wolferth and Geckeler 1943 Kent 1914) or through abnormal conducting fibres arising from the upper part of the bundle of His (Wolferth and Wood 1933) such as those described by Mahaim (1931) The passage of the excitatory impulse down such an alternative pathway might well account for premature right ventricular stimulation Experimental short circuits of the kind envisaged were devised by Butterworth and Poindexter (1942) the classical appearances of the Wolff Parkinson White syndrome resulted

The condition is congenital occurs in both sexes equally and is often unstable as shown by serial electrocardiograms indeed normal and abnormal complexes may alternate (fig 4 32b) On the whole normal conduction is encouraged by atropine abnormal conduction by cholinergic activity (Duthie 1946) The heart is otherwise normal in at least 70 per cent of cases and heart disease when present is probably coincidental Pre excitation is clinically and academically important on account of its association with paroxysmal tachycardia and is easily overlooked because casual electrocardiograms may be normal Paroxysmal tachycardia occurs in 50 to 60 per cent of cases (Willius and Carryer 1946) and is often closely related to effort Electrocardiograms obtained during attacks suggest that their mechanism may depend upon a circus movement the impulse travelling down the bundle of His and back through the short circuit (fig 4 33a) or down the short circuit and back through the bundle of His (fig 4 33b) the former being more common Both types of paroxysm may occur in the same patient as in the illustrations In this particular case the second type of paroxysm was provoked by mecholin and before normal rhythm was resumed there was a period of transition in which abnormal P waves appeared immediately after certain QRS complexes (fig 4 33c) causing a single premature ventricular beat and suggesting circus movement due to retrograde conduction through the bundle of Kent or similar structure initial excitation having passed through the bundle of His Similar P waves may be seen in the upper half of figure 4 33b but these fail to excite the ventricles Occasionally attacks resemble auricular flutter or fibrillation and the ventricular rate may be exceptionally fast A case described by Littmann and Tarnower (1946) had an irregular ventricular rate of 340 per minute

AURICULAR FLUTTER

Physiology Auricular flutter in man was so named by Jolly and Ritchie (1910) after obtaining the first electrocardiographic records of the condition and was attributed to a circus movement by Lewis (1918-20) The

excitatory impulse was believed to travel round a ring of auricular tissue, such as the mouths of the ventricle as proved possible by the physiological researches of Mines (1913) Rosenblueth and Ramo (1947) apparently confirm Lewis' views. Using a high speed cinematograph technique however Prinzmetal (1950) has disproved this thesis and has shown that auricular flutter and fibrillation like auricular ectopic beats and paroxysmal auricular tachycardia depend upon the presence and behaviour of an irritable focus in auricular muscle. The speed of the auricular beats ranges between 260 and 340 per minute and its rhythm is regular. As the A-V node can rarely transmit impulses faster than 210 to 220 per minute, physiological heart block results the ventricles usually responding to every second impulse. If the auricular rate is slower however and approaches 200 per minute as it may under the influence of quinidine the ventricles may be able to keep pace (Lewis 1925). If vagal tone is increased as by carotid sinus pressure a greater degree of physiological block results and an A-V ratio of 4:1 or so may be established. Sometimes the ventricular response is irregular.

Clinical features incidence and etiology. Clinically flutter should be suspected in any patient presenting a regular heart rate of 120 to 170 per minute uninfluenced by effort emotion or change of posture whether there are other indications of heart disease or not. The first heart sound often varies in intensity according to the time relationship between auricular and ventricular contractions (Harvey and Levine 1948).

Flutter is a relatively uncommon but capricious rhythm and may occur when least expected. It is twice as common in men as in women and its incidence increases with age being rare under 30 and most frequent (88 per cent) between the ages of 40 and 70. It is very rare in otherwise normal individuals it may complicate such diverse conditions as meningitis pneumonia cholecystitis or carcinoma of the colon. In 90 per cent of cases however it is associated with organic heart disease especially rheumatic hypertensive ischaemic or pulmonary and may then precipitate or complicate congestive heart failure. According to Campbell (1947) angina pectoris develops in 25 per cent of paroxysms. Attacks are commonly transient and have the same abrupt onset as paroxysmal tachycardia but they tend to last longer, being measured in weeks rather than hours and may occasionally persist for years. Lewis (1937) described a case in a parson which had continued for 24 years.

Diagnosis is facilitated by carotid sinus pressure, which often causes abrupt temporary slowing of the ventricular rate as described previously whereas in sinus tachycardia slowing is commonly slight and in paroxysmal tachycardia it is usually absent unless the attack is terminated. Electrocardiography is advised in all suspected cases however and reveals a continuous series of rapid regular auricular f waves (fig 4.34a) without intervening isoelectric periods. When there is 2:1 block, one f wave is more or less obscured by the QRS complex so that the nature of the

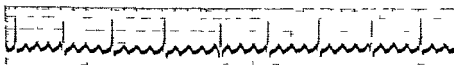


Fig. 4 34 (a)—Auricular flutter with 4-1 A V block

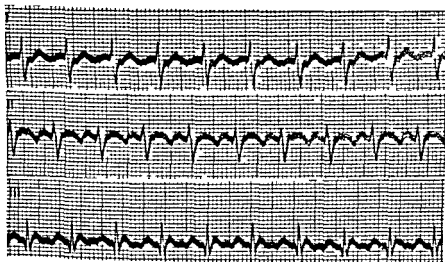


Fig. 4 34 (b)—Auricular flutter with 2-1 A V block

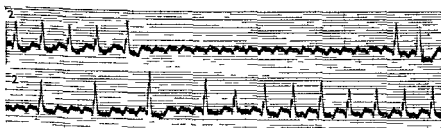


Fig. 4 34 (c)—Auricular flutter clarified by means of carotid sinus compression

tachycardia may remain uncertain (fig 4 34b). Carotid sinus pressure aids analysis by increasing the degree of block and so unmasking such hidden f waves (fig 4 34c).

Treatment The patient should be put to bed and treated with adequate doses of digitalis beginning with 9 grains (0.6 G) of the powdered leaf followed by 6 grains (0.4 G) and then by 3 grains (0.2 G) at six hourly intervals and continuing with 2 grains (0.13 G) t d s until serial electrocardiograms show that auricular fibrillation has been established. The drug

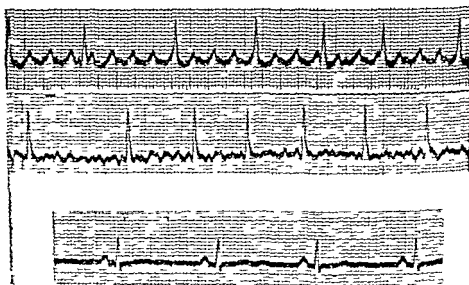


Fig 4 35—Auricular flutter treated with digitalis. Auricular fibrillation is induced but on withholding the drug normal rhythm is resumed.

is then withheld in the hope that normal rhythm may be resumed spontaneously (fig 4 35). Electrocardiographic control is necessary because the slow irregular ventricular response which results from such doses of digitalis is no proof of auricular fibrillation under the circumstances. Adequate supervision is important owing to the heavy dose of digitalis usually required to induce fibrillation and if toxic symptoms appear dangerous before this result is achieved the attempt may have to be abandoned. The effect of digitalis is twofold as already indicated: it encourages the irritable focus to assume the properties associated with auricular fibrillation and by depressing conduction in the bundle of His it slows the ventricular rate. It was hitherto believed that normal rhythm was resumed when the circus movement was broken by the head of the wave meeting a refractory tail (Lewis 1925) for circus movement could not occur unless there was a gap of responsive tissue just ahead of the wave. Digitalis either during its administration or when it was suspended was thought to close the gap by having an unequal and favourable effect on conduction and on the

refractory period Obviously if conduction were quickened and the refractory period prolonged in auricular tissue the hypothetical gap would close Naturally no drug has this effect those quickening conduction also shortening the refractory period (like the cholinergic bodies) and *vice versa* (like quinidine) The action of digitalis is complicated by its cholinergic effect the *f* waves are never retarded but they may be accelerated especially in those cases which are made to fibrillate (Wedd 1924)

Quinidine should not be given alone to cases of flutter in the first instance for by depressing the irritable focus and slowing the auricular rate it may allow the ventricles to keep pace rapid tachycardia resulting When auricular fibrillation has been established however the resumption of normal rhythm may be encouraged by quinidine in doses of 5 to 10 grains (0.32 to 0.65 G) two hourly to a maximum of 40 to 45 grains (2.5 to 3 G) in one day or it may be given to resistant cases of flutter so long as the ventricular response is blocked by digitalis

If flutter continues despite all efforts to break it the patient should be kept on a maintenance dose of digitalis sufficient to control the ventricular rate but the result is rarely satisfactory for short of digitalis intoxication tachycardia due to 2:1 ventricular response is apt to develop on little provocation

In all cases attention should be paid to any associated disease cardiac or otherwise and to combating congestive heart failure

AURICULAR FIBRILLATION

Physiology According to Prinzmetal (1930) two types of auricular contractions may be seen by means of a high speed cinematograph in experimental auricular fibrillation (1) minute irregular contractions which he has called M contractions involving a small area of auricular wall (0.03 x 3 mm) and (2) large rhythmic wave like contractions (L contractions) which sweep across the auricle 400 to 600 times per minute without pursuing a circus pathway Blocking a hypothetical circuit round the mouths of the venæ cavæ had no effect on these waves Direct auricular leads recorded by means of a cathode ray oscillograph showed very small M waves at 10 000 to 40 000 per minute and large *f* waves corresponding to the L contractions The M waves did not occur in flutter Lewis's theory of circus movement appears to be untenable

At speeds of 320 to 380 electrocardiograms from chest leads placed over the right auricle show *f* waves which at times are regular and even as in flutter and which at other times are irregular and uneven as in fibrillation (fig 4.36) At faster rates the *f* waves are always irregular in time and shape and the ventricular response is commonly rapid and chaotic varying between 100 and 200 per minute (fig 4.37) Sometimes and of course in treated cases when there is partial auriculo-ventricular block the ventricular rate is relatively slow Occasionally there is complete

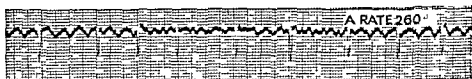


Fig 4 36—Lead CR1 showing coarse auricular fibrillation or impure flutter

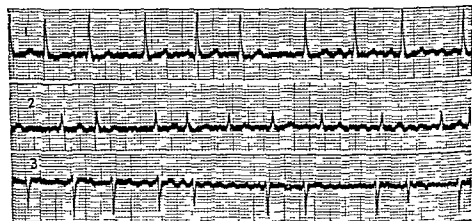


Fig 4 37—Auricular fibrillation

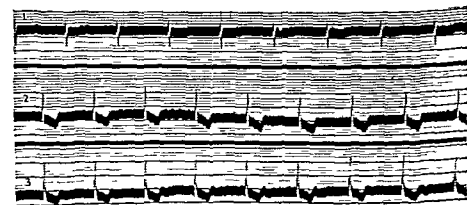


Fig 4 38—Auricular fibrillation with complete A V dissociation due to digitalis

heart block usually in cases treated with digitalis over a long period of time and the ventricular rate is not only slow but regular (fig 4 38)

Etiology Auricular fibrillation is characteristically associated with mitral stenosis and toxic nodular goitre and is usually permanent with the former and paroxysmal with the latter. It is not uncommon however in the later stages of hypertensive and ischaemic heart disease. On the other hand it is rare in congenital heart disease in bacterial endocarditis (2 per cent) in any form of active carditis in young people in aortic valve disease (unless there is stenosis of the coronary ostia) in pulmonary heart disease in the high output group (apart from thyrotoxicosis) and in pericarditis (although it occurs in 33 per cent of cases of Pick's disease). Like flutter more over auricular fibrillation may occur in patients with no other evidence of heart disease it may complicate head injuries meningitis pneumonia and other infections in rare instances and it may even be found in apparently healthy persons. The most important single factor determining the incidence of auricular fibrillation in those diseases that favour its occurrence is the advancing age of the patient.

Clinical features Symptoms may be absent or negligible or the patient may complain of palpitations. If the ventricular rate is very rapid syncope or angina pectoris may result as with flutter and paroxysmal tachycardia. The mechanical inefficiency and nutritional hazards resulting from the rapid irregularity of the heart beat often lead to congestive failure when there is underlying heart disease on the other hand auricular fibrillation may be precipitated by congestive failure from other causes.

Diagnosis The clinical diagnosis rests upon the recognition of a chaotic cardiac rhythm i.e. one without any semblance of order and must be distinguished from sinus arrhythmia from ectopic beats and from auricular flutter. Sinus arrhythmia should be recognised by its relation to respiration and ectopic beats by the perception of some fundamental order but multiple auricular ectopic beats especially when associated with sinus arrhythmia and auricular flutter with an irregular ventricular response may be most confusing. Electrocardiography is therefore advised in all suspected cases.

Treatment All cases in which the ventricular rate is accelerated should be treated with digitalis. When there is no urgency a simple and safe method is to give powdered digitalis leaf 3 grains (0.2 G) t.d.s. on the first day 2 grains (0.13 G) t.d.s. on the second and 1 grain (65 mg) t.d.s. thereafter until the ventricular rate is controlled. Subsequently a maintenance dose of 1 grain (65 mg) b.i.d. is usually sufficient. When a quicker effect is desired the method described for cases of auricular flutter is advised. In urgent cases with very rapid ventricular rates and severe congestive heart failure digoxin by the intravenous route may be preferable but is not without danger and should never be given in full doses to any patient who may have had digitalis within the previous six weeks or who still shows a digitalis effect in the electrocardiogram. The initial maximum dose is 1.5

mg and this may be followed by 0.5 mg and then by 0.25 mg, at intervals of not less than two hours and not more than six hours. In favourable circumstances the ventricular rate may be controlled within half an hour; an oral maintenance dose should then replace the later intravenous doses just recommended. Strophanthin may be used instead of digoxin as Ouabain; it may be given in an initial dose of 1.0 mg intravenously, followed by 0.5 mg and then by 0.25 mg six hourly until the desired effect is obtained. As strophanthin is all excreted within forty-eight hours, it is preferable to digoxin when a cumulative effect is not desired.

Other preparations of digitalis may be given by mouth, the dose being calculated according to the following table of equivalent strengths:

Powdered digitalis leaf	1 grain (65 mg)
Tincture of digitalis	10 minims (0.6 ml)
Digoxin	0.25 mg
Digitoxin (Nativelle's Digitaline)	0.075 mg

The practitioner is advised to become thoroughly familiar with a few reliable preparations. Digoxin and digitoxin have the advantage of being pure crystalloids of fixed potency. Digoxin appears to be excreted more quickly than digitoxin. The tincture loses strength with the passage of time and when mixed with other drugs and is therefore least reliable. The powdered leaf has been the standard preparation in this country for many years but is being gradually displaced by digoxin. A point of importance when calculating equivalent doses of different preparations is that the maximum single dose of the more rapidly excreted drugs such as strophanthin and digoxin is smaller than would be assessed by the table of equivalent strengths; thus it is safe to give an average adult 9 grains (0.6 G) of the powdered leaf as a massive single dose and to follow it at six hourly intervals by 6 grains (0.4 G) and then by 3 grains (0.2 G) but it is not safe to give 2.25 mg of digoxin followed by 1.5 mg and then by 0.75 mg although its maintenance dose is 0.25 mg twice daily. Massive oral doses of digoxin should not exceed 1.5 mg, 1 mg and 0.5 mg at six hourly intervals.

Toxic symptoms include anorexia, nausea, vomiting, diarrhoea, ectopic beats, nodal rhythm, heart block, paroxysmal tachycardia and sudden death from ventricular fibrillation. Nausea and coupling due to ectopic beats are the best indications that the accumulated dose of digitalis is approaching dangerous concentration. Unfortunately the worse the heart the closer the therapeutic dose becomes to the toxic; the margin is never great. The vagal effects may be relieved by atropine.

The correct maintenance dose must be worked out for each individual receiving the drug but it averages 0.5 mg of digoxin daily, ranging between 0.25 and 0.75 mg.

Attempts to restore normal rhythm with quinidine should be made in all cases in which there is no evidence of intrinsic heart disease and especially

in cases of successfully treated thyrotoxicosis and perhaps when auricular fibrillation is thought to have occurred prematurely or unexpectedly having been precipitated by some passing infection such as tonsillitis or pneumonia or by some other factor which either no longer operates such as pregnancy or which is itself controllable such as dental sepsis. When fibrillation develops in the natural course of heart disease however e.g. in mitral stenosis or hypertensive heart disease attempts to restore normal rhythm end in immediate or remote failure and should therefore be avoided as the procedure is not without risk.

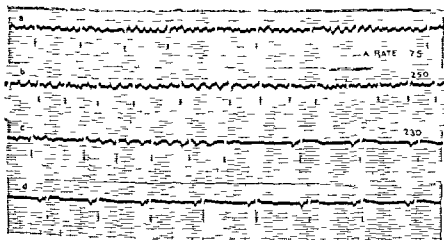


Fig 4.39—Auricular fibrillation treated with quinidine. The f waves slow down from 375 to 230 per minute before normal rhythm is restored.

Quinidine should be given by mouth in doses of 5 grains (0.32 G) two hourly on the first day followed by 10 grains (0.65 G) two hourly on the second and by 15 grains (1 G) two hourly on the third to a maximum of 40 to 45 grains (2.6 to 2.9 G) per day the course being terminated immediately the rhythm returns to normal. A maintenance dose of 5 grains (0.32 G) t.d.s. is continued for a month in successful cases.

Quinidine depresses the activity of the irritable focus retarding its periodicity and often abolishing it altogether (about 50 per cent of cases). As the f waves slow down (fig 4.39) they may assume the regularity of flutter and if their speed approaches 200 per minute there is danger of a 1:1 ventricular response. Tachycardia so provoked by quinidine may be prevented by preliminary digitalis therapy and a maintenance dose of digitalis is advised throughout the quinidine course. The theoretical consideration that digitalis and quinidine have partly opposing actions does not prejudice successful practical results.

Other complications of quinidine therapy include hypersensitivity and embolism. Hypersensitivity may result in generalised oedema, urticaria, purpura, fever, vomiting and collapse although such symptoms are rare.

it is customary to give an initial trial dose of 3 grains (0.2 G). Important systemic emboli occur in about 5 per cent of all cases in which normal rhythm is restored and are due to the expulsion of left auricular thrombi. The risk is more or less proportional to the length of the period of fibrillation and is greatly increased by congestive heart failure. Intracardiac thrombi are rare in thyrotoxic heart disease even under the most unfavourable circumstances owing to the rapid circulation associated with it. Pulmonary emboli also occur but rarely prove troublesome.

VENTRICULAR FIBRILLATION

Faradic stimulation of the ventricles invariably induces incoordinated fibrillation of the muscle which usually persists after cessation of the exciting cause. The heart muscle is unable to expel its contents and syncope

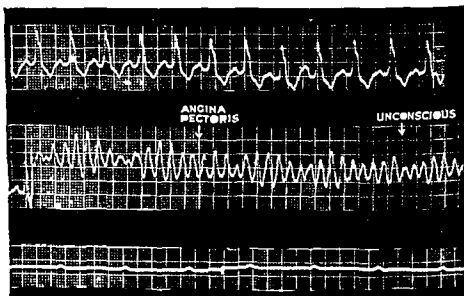


Fig. 4.40.—Ventricular fibrillation causing sudden death in a case of ischaemic heart disease

occurs abruptly. Spontaneous recovery may occur especially in young healthy animals but sudden death is the rule. When the heart is unduly excitable as in asphyxia digital pressure or gently scratching the surface of the ventricle with a pin may be sufficient to induce ventricular fibrillation (MacWilliam 1887). Certain drugs may initiate the phenomenon notably adrenaline, chloroform and digitalis. Coronary occlusion is also known to be an exciting cause.

Clinically ventricular fibrillation is often responsible for sudden death especially in ischaemic heart disease (fig. 4.40), aortic stenosis, syphilitic aortic incompetence, diphtheritic carditis and complete heart block. It also

explains sudden death following intravenous injections of digitalis mercurial diuretics adrenaline and other drugs

Treatment is of little avail The intracardiac injection of quinidine sulphate 3 to 5 grains (0.2 to 0.32 G) or of 5 to 10 ml of 1 to 2 per cent procaine may be tried if circumstances are favourable Quinidine may also be used as a prophylactic agent when the risk of ventricular fibrillation is great e.g. following coronary thrombosis or in status anginosus

REFERENCES

- Barker P S MacLeod A G and Alexander J (1930) The excitatory process observed in the exposed human heart *Amer Heart J* 5 720
- Bayley R H (1934) The frequency and significance of right bundle branch block *Amer J med Sc* 188 236
- Bouveret L (1889) Concerning essential paroxysmal tachycardia *Ret de Méd* 9 753
- Braun Menendez E and Solari L A (1939) Ventricular asynchronism in bundle branch block *Arch intern Med* 63 830
- Butterworth J S and Poindexter C A (1942) Short PR interval associated with prolonged QRS complex clinical and experimental study *Ibid* 69 437
- Campbell M (1944) Complete heart block *Brit Heart J* 6 69 — (1945) Paroxysmal tachycardia and 2:1 heart block *Ibid* 7 183 — (1947) The paroxysmal tachycardias *Lancet* ii 681
- Castleden L I M (1941) The effect of potassium salts on cardiac irregularities *Brit med J* i 7
- Cotton R P (1867) Notes and observations upon a case of unusually rapid action of the heart (232 per minute) *Ibid* i 629
- Duthie R J (1946) Mechanism of the Wolff Parkinson White syndrome *Brit Heart J* 8 96
- Evans W (1944) The unity of paroxysmal tachycardia and auricular flutter *Ibid* 6 221
- Formigne P (1938) Apnoea or convulsions following standstill of the heart *Amer Heart J* 15 19
- Graybiel A and White P D (1936) Complete auriculo ventricular dissociation A clinical study of seventy two cases with a note on a curious form of auricular arrhythmia frequently observed *Amer J med Sc* 192 334
- Griffith T W (1921) A clinical study of three cases of heart block *Brit med J* i 763
- Gross L (1921) The blood supply to the heart New York
- Harvey W P and Levine S A (1948) The changing intensity of the first heart sound in auricular flutter an aid to the diagnosis by auscultation *Amer Heart J* 35 924
- Hay J (1906) Bradycardia and cardiac arrhythmia produced by depression of certain functions of the heart *Lancet* i 139
- Holzmann M and Scheff D (193) Über Elektrokardiogramme mit verkürzter Vorhof Kammer Distanz und positiven P Zacken *Ztschr f klin Med* 121 404
- Jolly W A and Ritchie W R (1910) Auricular flutter and fibrillation *Heart* 2 177

- Kay H B (1948) Ventricular complexes in heart block *Brit Heart J* 10 177
- Keith A and Flack M (1907) The form and nature of the muscular connexions between the primary divisions of the vertebrate heart *J of Anat and Physiol* 41 172
- Kent A F S (1914) Observations on the auriculo ventricular junction of the mammalian heart *Quart J exper Physiol* 7 193
- Kountz W B (1936) Revival of human hearts *Ann intern Med* 10 330
- Lawrence J S and Forbes G W (1944) Paroxysmal heart block and ventricular standstill *Brit Heart J* 6 53
- Levine S A (1948) Auscultation of the Heart *Ibid* 10 213
- Lewis T (1925) The mechanism and graphic registration of the heart beat London — (1937) Auricular flutter continuing for twenty four years *Brit med J* 1: 1248 — Feil H S and Stroud W D (1918-20) Observations upon flutter and fibrillation Part II The nature of auricular flutter *Heart* 7 191
- Littmann D and Tarnower H (1946) Wolff Parkinson White syndrome A clinical study with report of nine cases *Amer Heart J* 32 100
- McMichael J and Sharpey Schafer E P (1944) Cardiac output in man by a direct Fick method *Brit Heart J* 6 33
- McWilliam J A (1887) Fibrillar contraction of the heart *J Physiol* 8 296
- Mahaim I (1931) Les maladies organiques du faisceau de His Tawara Paris
- Major R H (1923) Stokes Adams disease due to gummata of the heart *Arch intern Med* 31 857 — (1931) Classic descriptions of disease Springfield Illinois U S A
- Mann H (1931) Interpretation of bundle branch block by means of monocardioqram *Amer Heart J* 6 447
- Mines G R (1913) On dynamic equilibrium in the heart *J Physiol* 46 349
- Parkinson J Papp F and Evans W (1941) The electrocardiogram of the Stokes Adams attack *Brit Heart J* 3 171
- Prinzmetal M et al (1950) Mechanism of the auricular arrhythmias *Circulation* 1 241
- Rosenblueth A and Ramos J G (1947) Studies on flutter and fibrillation II The influence of artificial obstacles on experimental auricular flutter *Amer Heart J* 33 677
- Sampson J J and Anderson E M (1932) The treatment of certain cardiac arrhythmias with potassium salts *J Amer med Ass* 99 2257
- Spens T (1793) History of a case in which there took place a remarkable slowness of the pulse *Medical Commentaries (Edinburgh)* 7 463
- Stokes W (1846) Observations on some cases of a permanently slow pulse *Dublin quart J med Sc* 2 73
- Szekely P (1946) The action of Magnesium on the heart *Brit Heart J* 8 115
- Tawara S (1906) Das Reizleitungssystem des Säugetierherzens Jena
- Tung C L and Cheer S N (1933) A correlation of clinical and electrocardiographic findings in human bundle branch block *Chinese med J* 47 15
- Wedd A M (19-4) Notes on the action of certain drugs in clinical flutter *Heart* 11 87
- Wenckebach K F (1899) Zur Analyse des unregelmässigen pulses II Ueber den regelmässig intermittierenden Puls *Zetschr f klin Med* 37 475
- White P D (1915) A study of atrio-ventricular rhythm following auricular flutter *Arch intern Med* 16 517

Willius F A and Carryer H V (1946) Electrocardiograms displaying short P R intervals with prolonged QRS complexes: an analysis of 65 cases. *Proc May Clin* 21 438

— Johnston F D Hill I G W MacLeod A G and Barker P S (1934) The significance of electrocardiograms characterised by an abnormally long QRS interval and by broad S deflections in lead 1. *Ibid* 9 459

Wilson F N MacLeod A G and Barker P S (1932) The order of ventricular excitation in human bundle branch block. *Amer Heart J* 7 305

Wolferth C C and Margolies A (1935) Asynchronism in contraction of the ventricles in the so called common type of bundle branch block: its bearing on the determination of the side of the significant lesion and on the mechanism of split first and second heart sounds. *Ibid* 10 425

— and Wood F C (1933) The mechanism of production of short P R intervals and prolonged QRS complexes in patients with presumably undamaged hearts: hypothesis of an accessory pathway of auriculo ventricular conduction (Bundle of Kent). *Ibid* 8 297

Wolff L Iarkinson J and White P D (1930) Bundle branch block with short P R interval in healthy young people prone to paroxysmal tachycardia. *Ibid* 5 685

Wood F C Jeffers W A and Wolferth C C (1935) Follow up study of sixty four patients with right bundle branch conduction defect. *Ibid* 10 1056

— Wolferth C C and Geckeler G D (1943) Histologic demonstration of accessory muscular connexions between auricle and ventricle in a case of short P R interval and prolonged QRS complex. *Ibid* 25 454

Yater W M (1938) Pathogenesis of bundle branch block: review of the literature: report of sixteen cases with necropsy and of six cases with detailed histologic study of the conduction system. *Arch intern Med* 62 1

CHAPTER V

HEART FAILURE

HEART failure has been defined as a condition in which the heart fails to discharge its contents adequately (Lewis 1933). The words may be applied logically to the heart as a whole or to one or other ventricle. The adjective congestive is often added and has come to mean heart failure with systemic congestion i.e. with elevation of the systemic venous pressure and engorgement of the liver, with or without dropsy. Right ventricular failure has a similar meaning but implies also that the left ventricle is relatively healthy. Left ventricular failure is characterised by congestion of the lungs only.

MECHANISM

The mechanism and even the definition of heart failure have been debated for over a century and are still a source of controversy. The back pressure theory so well expressed by James Hope in 1832 which incorporates the idea of independent ventricular failure maintains that when a ventricle fails to discharge its contents adequately blood accumulates behind it and the pressure rises in the respective auricle and venous system. After holding sway for nearly a century this conception was replaced by the forward failure hypothesis of Mackenzie (1913) who believed that congestion depended upon failure of sufficient propulsion from behind and who insisted that the heart failed as a whole. Before the second world war opinion reverted sharply to Hope's view the arguments in its favour being well marshalled by Harrison (1935) and by Fishberg (1939) but the newer methods of investigation which provided much of the data upon which these arguments were based were crude and subsequent technical refinements have disproved many of them. The modern attitude has been shaped largely by the work of Cournand in the U.S.A. and of McMichael and Sharpey Schafer in England.

By means of intracardiac catheterisation these investigators have provided reliable data concerning the cardiac output and the pressures in the right side of the heart and McMichael and Sharpey Schafer have developed a theory of congestive failure which is in accordance with Starling's Law of the heart (McMichael 1947). According to this law (Starling 1918) the cardiac output is proportional to the venous filling pressure (right auricular pressure minus the negative intrathoracic pressure) until a critical level is reached after which any further rise of venous pressure results in overloading and a fall in output (fig. 5.01). It is known that in patients with

severe anemia a raised venous pressure is primarily a physiological adjustment which serves to maintain a high cardiac output and that artificial alterations of the venous pressure results in changes of cardiac output in harmony with Starling's Law (Sharpey Schafer 1944) Should a further rise of venous pressure result in a lower cardiac output the heart is said to be overloaded

McMichael and Sharpey-Schafer (1944) after making further observations on the relationship between the right auricular pressure and the

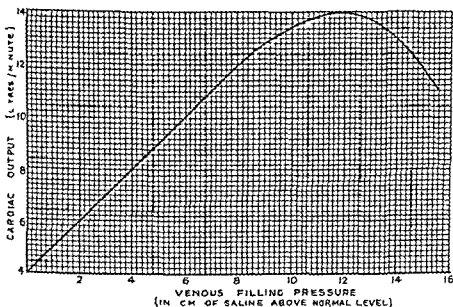


Fig 501—Relationship of cardiac output to venous filling pressure (Starling's curve)

cardiac output in health and disease put forward the hypothesis that elevation of the venous pressure is primary in all forms of congestive heart failure and may be regarded as a compensatory mechanism designed to increase the cardiac output. For a time and particularly in certain hyperkinetic circulatory states (e.g. anemia, arterio-venous aneurysm) in which the cardiac output is high the mechanism is successful but sooner or later overloading occurs. According to this hypothesis congestive heart failure might be defined as a state in which the heart can only maintain the requisite output by means of abnormal elevation of the venous pressure or as a state in which such elevation of the venous pressure has already resulted in overloading. The second clause defines a more advanced condition than does the first and it may well be objected that the first does not in fact define congestive failure at all but a compensated state which has no right to the title. Congestive heart failure is a clinical syndrome characterised by elevation of the venous pressure and distension of the liver with

or without dependent œdema the result of some cardiac fault. If these manifestations can be demonstrated when the heart is not yet overloaded the definition must be allowed.

The clinical facts are these. In anæmia arterio venous aneurysm and extensive active Paget's disease of bone elevation of the venous pressure without demonstrable distension of the liver is commonly associated with a high cardiac output. If the venous pressure is raised further the output usually rises; if it is reduced the output usually falls. Such behaviour has been labelled 'high output failure' and is covered by the first of the two definitions given above. Cervical venous pulsation is rarely visible above clavicular level when the patient reclines at an angle of 45 degrees; in other words it is rarely more than 3 or 4 cm. above the sternal angle in this position. When the venous pressure is raised more conspicuously the liver becomes palpable and the heart is probably overloaded. In thyrotoxicosis high cardiac outputs are maintained chiefly by means of tachycardia. Clinical elevation of the jugular venous pressure is associated with hepatic enlargement and is only seen when the heart is overloaded. The cardiac output is then usually low. In pulmonary heart disease secondary to emphysema elevation of the venous pressure is commonly associated with hepatic engorgement and with a moderate increase in cardiac output. The latter usually falls when the venous pressure is further raised, but it also falls when the venous pressure is lowered. In most other forms of heart disease elevation of the venous pressure is associated with hepatic engorgement and with a low cardiac output; further elevation of the venous pressure results in further reduction of the cardiac output.

Clinicians are therefore likely to favour the view that congestive heart failure is a state in which the heart is overloaded, i.e. a state in which further elevation of the venous pressure causes a reduction in cardiac output. The best clinical indication of this may be demonstrable distension of the liver. Whatever the final verdict on this vexed question, it is certainly true to say that in congestive heart failure the venous pressure rises primarily behind the chamber chiefly involved, i.e. in the left auricle and pulmonary veins in mitral stenosis and left ventricular failure; in the right auricle and systemic veins in tricuspid valve disease and right ventricular failure. Moreover in mitral stenosis and left ventricular failure secondary elevation of the systemic venous pressure is the rule sooner or later, whether this be regarded as evidence of right ventricular failure or otherwise.

CAUSES OF HEART FAILURE

The heart may fail because it is overburdened by a raised ventricular pressure or by a raised cardiac output, or because the health of the myocardium is impaired by inadequate or faulty nutrition, metabolic disorder, intoxication, or intrinsic disease. High outputs are tolerated better than high pressures, but myocardial ill health is probably even more important.

Contributory factors include physical effort obesity anxiety mental stress disturbances of rate or rhythm infection fever extremes of temperature and pregnancy but all these are better expressed in more fundamental terms for example infection may increase the cardiac output and impair the health of the myocardium anxiety and cold may raise the blood pressure and so forth

Viewing the subject in this way it should be clear that a high cardiac output is no more incompatible with heart failure than is hypertension that a heart capable of pumping ten litres of blood per minute is not necessarily better than one capable of maintaining a diastolic blood pressure of 140 mm. of Hg Each is a measure of part of the total cardiac work performed neither alone is a sufficient measure of cardiac efficiency although their behaviour under certain experimental conditions may be Moreover the signs and symptoms of heart failure are largely due to alterations of pressure and volume in the pulmonary or systemic venous systems In left ventricular failure for example the redistribution of volume is the result of a short lived discrepancy between left and right ventricular outputs Although the balance must be restored quickly the consequences cannot be rectified until the process is reversed It should again be clear that such disturbances cannot be detected by casual estimations of the right ventricular output

LEFT VENTRICULAR FAILURE

When the left ventricle fails to discharge its contents adequately blood accumulates in the pulmonary circulation and the pressure rises in the left auricle and pulmonary veins

ETIOLOGY

Left ventricular failure may result from any disease which imposes an undue burden on the left ventricle or which interferes with its health These diseases include systemic hypertension from any cause aortic valve disease and myocardial infarction In systemic hypertension the left ventricle may fail either because it is unable to meet the stress imposed upon it or because it is enlarged so greatly that it cannot obtain sufficient nourishment As the nutritional demands of an individual muscle fibre depend upon its cubic volume and the nutritional supply is limited by its surface area there is an increasing discrepancy between the two as the muscle enlarges which sooner or later becomes critical (Gross and Spark 1937) In acute nephritis and malignant hypertension a rapid rise of blood pressure may cause left ventricular failure before there has been appreciable hypertrophy of muscle on the other hand in long standing cases of essential hypertension with gross enlargement of the left ventricle failure may occur even though the blood pressure has fallen to within normal limits failure then being attributed to nutritional breakdown In aortic valve disease in addition to these two factors there may be further interference

with nutrition as a result of poor coronary filling due to a low mean blood pressure in aortic stenosis and to obstruction of the mouths of the coronary vessels in syphilitic aortic incompetence. The cause of failure in uncomplicated ischæmic heart disease with myocardial infarction is due entirely to interference with ventricular nutrition resulting from coronary occlusion.

CLINICAL FEATURES

The symptoms of left ventricular failure are undue breathlessness on effort, orthopnoea, paroxysmal cardiac dyspnoea and acute pulmonary oedema. The findings include bilateral basal pulmonary râles, diminished vital capacity and lung volume, radiological evidence of pulmonary congestion and hydrothorax and prolongation of the pulmonary circulation time. The diagnosis is supported by gallop rhythm, pulsus alternans and Cheyne Stokes breathing and is confirmed by the demonstration of a suitable cardiovascular disease, e.g. systemic hypertension, aortic valve disease, or myocardial infarction.

Undue breathlessness on effort. Breathlessness on effort is physiological. When a patient complains of breathlessness, he means that he is winded by physical work that did not distress him previously. The symptom, *per se*, does not of course necessarily indicate heart disease. Other common causes including psychoneurosis, obesity, chronic bronchitis, asthma, emphysema and anæmia. Breathlessness due to left ventricular failure depends upon pulmonary congestion, which both reduces the vital capacity and reflexly stimulates respiration.

Orthopnoea, paroxysmal cardiac dyspnoea and pulmonary oedema. As these three conditions depend on variations of the same fundamental mechanism, they are considered together. When a patient adopts the upright or sitting position in order to breathe comfortably, he may be said to have orthopnoea. Although an almost constant sign of left ventricular failure, it is by no means pathognomonic for it may be found in severe mitral stenosis, bronchial asthma and in pericardial effusion. The vital capacity is reduced in all these conditions and is greater in the upright than in the horizontal position, but its relationship to orthopnoea is not necessarily direct. More over, its increase in the erect position is greater than can be explained by descent of the diaphragm. The discrepancy is due to concomitant changes in the pulmonary circulation, the amount of blood in the lungs being greater perhaps by as much as 500 ml. in the horizontal than in the erect position (McMichael 1939). The redistribution of blood depends upon the geographical relationship of the auricles to their respective venous systems. As the right auricle is nearer the head than the feet, the pressure within it rises when the body is tilted head up, owing to the influence of gravity. The right ventricle responds according to Starling's Law and pumps more blood into the lungs in the horizontal than in the vertical position. The pressure within the left auricle, however, which is situated more or less in the centre of the lungs, is not directly influenced by gravity and the left

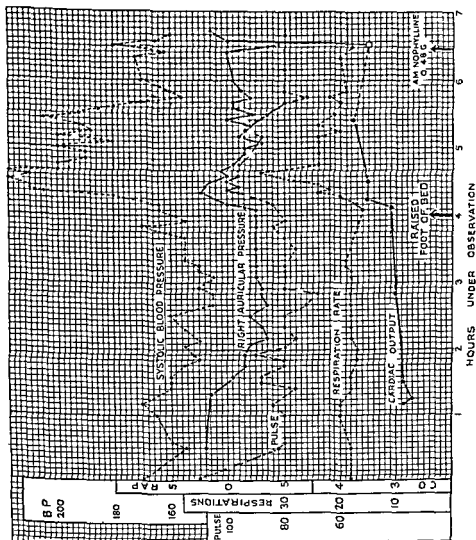


Fig 502—Graph illustrating typical changes in blood pressure, right auricular pressure, pulse rate, respiration rate, and cardiac output in an attack of paroxysmal cardiac dyspnoea initiated in a patient with left ventricular failure by raising the foot of the bed. The effect of aminophylline is shown at the end.

ventricular output does not therefore, immediately keep pace with that of the right. Only when the left auricular pressure rises proportionately owing to an increased volume of blood in the pulmonary venous system, will the balance be restored. As patients with left ventricular failure already have pulmonary congestion the extra engorgement which results from adopting the horizontal position may prove critical and may excite respiratory reflexes which provoke dyspnoea.

Paroxysmal cardiac dyspnoea usually occurs at night. The patient awakes with a feeling of suffocation and sits bolt upright gasping for breath, he may climb out of bed and open a window or walk about in an agitated way. In cases of simple orthopnoea this behaviour brings immediate relief but in paroxysmal cardiac dyspnoea the feeling of suffocation increases and the struggle for breath lasts for ten to twenty minutes. Coughing and wheezing are commonly associated (*cardiac asthma*) and the patient may complain of palpitations, faintness, or substernal tightness. The skin is pale, cyanosed and cold indicating profound vasoconstriction and sweating may be profuse. The blood pressure and venous pressure are both raised. Attacks usually subside spontaneously but may be repeated nightly or at intervals of days or weeks. In more severe cases pulmonary oedema develops. Widespread crepitations may be heard over the lungs and quantities of frothy pink or white watery fluid are expectorated.

Such attacks may sometimes be provoked by effort or by a rigor. They are easily induced experimentally in susceptible subjects by raising either the venous pressure or the blood pressure by artificial means (fig 5.02). The mechanism probably depends upon acute discrepancy between right and left ventricular outputs so that both the pressure and volume of blood in the pulmonary circulation reach critical levels. Measurements of pressure changes by means of an indwelling cardiac catheter in spontaneous nocturnal attacks indicate that the venous pressure may rise before the blood pressure. When attacks are induced by raising the venous pressure the cardiac output may rise. Thus although the heart is said to be failing it may in fact be performing more work than usual both with respect to blood pressure and output. The laboured breathing may be due in part to the extra effort required to inflate and deflate a turgid lung, the intrapleural pressure showing greatly increased fluctuations (Heyer *et al.*, 1948).

Spontaneous termination of the attack may be due to reduction of right auricular pressure by adoption of the upright posture or possibly to overloading of the weaker right ventricle so that the left has a chance to recover and restore the *status quo*. The difference between paroxysmal cardiac dyspnoea and acute pulmonary oedema is chiefly one of degree transudation of fluid from the capillaries into the alveolar spaces occurring when the intravascular hydrostatic pressure is sufficiently high or when other factors influence the fluid balance in favour of the tissues.

Bilateral basal pulmonary rales and hydrothorax Basal rales diminished

air entry into the lower lobes and some impairment of the percussion note at the bases are usual in left ventricular failure. Bedford and Lovibond (1941) found that hydrothorax was a common complication of pulmonary congestion from left ventricular failure and that although often bilateral tended to be more marked on the left side. Its occurrence may depend upon the fact that the visceral pleura is drained by the pulmonary rather than by the bronchial veins (Miller 1937).

Reduction of the vital capacity and lung volume The vital capacity is reduced by an amount equivalent to the extra quantity of blood distending the pulmonary circulation; it is reduced much more if there is pulmonary oedema or a large hydrothorax as well and by a further few hundred ml according to the degree of cardiac enlargement. Readings of 1000 to 1500 ml are common and may be as low as 500 ml when there is pulmonary oedema or hydrothorax.

The lung volume is reduced proportionately; the residual air remaining unchanged. This at once distinguishes the condition from emphysema in which a low vital capacity is associated with a normal lung volume and increased residual air.

Radiological signs of pulmonary congestion

Although the râles revealed by auscultation are most pronounced at the most dependent parts of the lung, the increased opacity seen in skiagrams is hilar and is due to engorgement of the pulmonary vessels (fig 5.03). During attacks of acute pulmonary oedema a fleecy mottling spreads out from the hilum on both sides (fig 5.04). Hydrothorax may also be revealed by X rays perhaps when unsuspected clinically. Confirmatory evidence of left ventricular failure may be obtained by noting the size and shape of the heart shadow.

Prolongation of the pulmonary circulation time The normal arm to tongue circulation time as measured by decholin or saccharin (page 13) averages 13.5 seconds but ranges between 9 and 18 seconds. As the time taken by the substance to travel from the left ventricle to the tongue may be neg-



Fig 5.03—Pulmonary congestion in left ventricular failure (case of syphilitic aortic incompetence)



() From 1 ft nit sala f har



(1) From mitral stenosis

1 g 504—A ute pul non ry ced ma

lected and as the journey from the antecubital vein to the right auricle takes only two or three seconds (Blumgart and Weiss 1927) the total arm to tongue time is governed chiefly by passage through the lungs. Using radium C intravenously, which can be detected at any given point in the circulation by means of a special radio sensitive instrument Blumgart and Weiss also showed that when the systemic venous pressure is raised in congestive heart failure the delay between the antecubital vein and the right auricle does not exceed five seconds even in gross cases. It follows that with pure right ventricular failure the arm to tongue circulation time should not exceed 23 seconds and should often be within normal limits in fact this is so. On the other hand in left ventricular failure the average time is 30 seconds (Wood 1936) and may be much longer. The delay is due to pulmonary congestion and occurs presumably on the venous side.

The arm to lung time The arm to lung time as measured by ether or amyl acetate (page 13) is said to be helpful in distinguishing primary left from pure right ventricular failure if the total arm to tongue time is also known. When the delay is proximal to the heart as in pure right ventricular failure the arm to lung time is delayed as much as the arm to tongue time on the other hand if there is further delay in the pulmonary veins as in primary left ventricular failure the arm to tongue time is disproportionately prolonged. Although theoretically this test might seem helpful in fact it is rarely so for two reasons first because the end point in the lung both with ether and amyl acetate is often unreliable and indefinite and second because it is easier and no less accurate to allow 1 to 5 seconds for delay proximal to the heart according to the degree of systemic venous engorgement.

GALLOP RHYTHM

When the rhythm of the heart sounds has three instead of two beats per musical measure or bar or when the metre of the heart beats has three instead of two syllables per poetical foot one may properly speak of triple rhythm. The term therefore covers all varieties of cadence in which three heart sounds are heard. Gallop rhythm on the other hand should have a stricter meaning and should be applied only to specified forms of triple rhythm as explained subsequently.

Mechanism Phonocardiography proves that there are really four normal heart sounds the auricular or presystolic sound (sometimes known as the fourth heart sound) associated with auricular systole and late ventricular distension the first heart sound due to mitral and tricuspid valve closure and to ventricular contraction the second heart sound due to closure of the aortic and pulmonary valves and the third heart sound which is attributed to sudden distension of the ventricles in the phase of rapid filling. Each of these sound is thus composed of at least two and at most four elements. Although these elements may not be strictly synchronous they are sufficiently so as a rule to produce but one obvious sound to the human ear.

On more careful analysis, however they may often be separated sufficiently to be detected individually by auscultation and we may then speak of split sounds. The word *split* describes the sound well and also indicates the mechanism of its production. The term *reduplication* is often used instead but has less to recommend it for it bears an accidental onomatopœtic resemblance to the sound of presystolic gallop and it is illogical to apply a word that means doubling to an act of division. Split sounds do not give the cadence of triple rhythm because of the close proximity of the separated elements.

The extra sound that is responsible for triple rhythm is usually an exaggerated auricular sound, the third heart sound or a summation of the two. Occasionally it is an additional systolic sound of unknown origin.

Presystolic gallop (auricular gallop) An audible auricular sound associated with a normal or slightly prolonged P-R interval gives rise to triple rhythm (presystolic or auricular gallop) with an amphibrachic metre (u ~ u). As it may be felt as well as heard it is best appreciated by means of a rigid

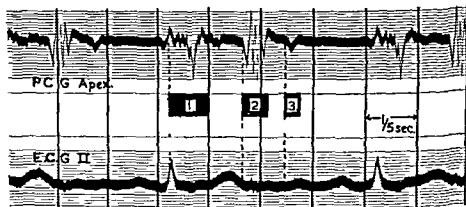


FIG. 505.—Phonocardiogram showing a normal third heart sound
(By G. I. J. D. F. D. H. Cote)

wooden stethoscope or with the naked ear so that tactile and aural senses may be allied. The presystolic sound is soft and dull and is usually localised to the region of the apex beat. In this situation it is pathognomonic of left ventricular stress. Occasionally it is heard best at the left border of the sternum when it may denote right ventricular stress. If the P-R interval is sufficiently prolonged the auricular sound may fall in mid or early diastole; if the heart rate is fast its true relation to the first or second heart sound cannot be determined clinically unless transient slowing is induced by means of carotid sinus compression. Presystolic gallop is never heard when there is auricular fibrillation.

Normal third heart sound When the extra sound occurs shortly after the second heart sound giving the metre of a dactyl (- u u) it may represent a normal or abnormal third heart sound (fig. 505). The normal third heart sound was well described by Gibson (1907). It is soft, low pitched and

usually accompanied by a palpable shock. It is more or less localised to the apex beat, varies in intensity with respiration and is accentuated when the subject lies on the left side especially if the venous pressure is raised by pressing on the abdomen. It may be heard in the great majority of children (but not in infants) in about 50 per cent of young adults occasionally in the middle aged and rarely in the elderly. Phonocardiography shows that the third heart sound synchronises with the latter half of the descending limb of the v wave of the jugular phlebogram and therefore with the period of rapid ventricular filling (Ohm 1913). It is attributed to sudden distension of the left ventricle at this time.

Protodiastolic gallop Abnormal third heart sounds are common in mitral stenosis constrictive pericarditis and in advanced hypertensive or ischaemic heart failure (protodiastolic gallop) especially when there is auricular fibrillation. The age and clinical condition of the patient emphasise their significance.

Summation gallop Summation of the auricular and third heart sounds can only occur when there is tachycardia or when the P R interval is sufficiently prolonged. With tachycardia the metre may seem to be anaæsthetic (u u -) dactylic (- u u) or amphibrachic (u - u) according to the fancy of the listener for the extra sound occurs in mid diastole. Summation sounds have no clinical significance if they disappear when the heart is slowed by carotid sinus compression (summation gallop) on the other hand such slowing may reveal an auricular sound or a normal or abnormal third heart sound.

Extra systolic sounds It is not uncommon for an extra sound to occur during ventricular systole. There are three varieties - the systolic click of left sided pneumothorax lesser systolic clicks possibly associated with pleuro pericardial adhesions and a third type in which the extra sound is dull and muffled and in no way like a click. Patients with partial left sided pneumothorax may complain of a loud clicking or bubbling noise synchronous with the heart beat. It may be so loud that it can be heard at a distance of several feet from the patient it varies markedly with respiration and with change of posture and is always transient. It is occasioned by the activities of bubbles of air between the heart and surrounding structures and only occurs when the pneumothorax is small so that clinically it is a late development appearing when most of the air has been absorbed (Scadding and Wood 1939). Lesser systolic clicks are heard from time to time in subjects who are perfectly well and according to Gallavardin (1913) may depend upon pleuro pericardial adhesions. In these cases the extra sound resembles a click but is not so impressive nor so variable as that associated with left sided pneumothorax. It may last for weeks months or years and may come and go without apparent reason. The third type (systolic gallop) is distinguished from greater and lesser systolic click by the character of the extra sound which is dull and muffled. Its mechanism is not yet understood. It is uncommon and when heard may be dis-

regarded for it occurs in apparently healthy persons. It should be distinguished from the widely split first sound of bundle branch block.

Note on nomenclature. Introduced by Professor Bouillaud, analysed and popularised by Potain (1876), the term gallop rhythm originally referred to that variety of triple rhythm which denoted impending or actual left ventricular failure, and in the presence of tachycardia is marvellously adapted to the sound it designates. But by 1900 Potain had extended the meaning of the *bruit de galop* to include presystolic, protodiastolic and systolic varieties, attributing these different metres to the same factors that are to-day held responsible. Thus historically it is not incorrect to regard gallop rhythm and triple rhythm as synonyms, but there is an advantage in excluding certain types of triple rhythm from the cadences embraced by the *bruit de galop*. Thus it is preferable and customary to speak of pre-systolic (auricular), protodiastolic, systolic and summation gallops on the one hand, and of systolic clicks, the normal third heart sound and the triple rhythm of mitral stenosis on the other.

PULSUS ALTERNANS

Pulsus alternans (Traube, 1872) is characterised by a regular rhythm in which the pulse beats are stronger and weaker alternately. It may be detected by palpation or more easily by sphygmomanometry, there being a difference of 5 to 20 mm. of mercury in the systolic pressure between

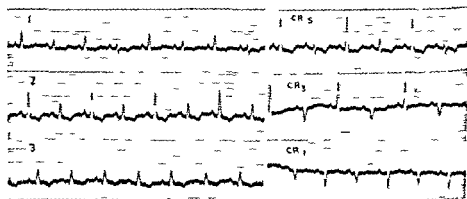


Fig. 506—Electrical alternation in a case of malignant disease in which the pericardium pulsus alternans was present.

alternate beats. It may be found in association with left ventricular failure, toxic carditis, paroxysmal tachycardia or auricular flutter. Clinically, alternation may be maintained as long as the heart is labouring, occasionally for as long as two or three years. Latent alternation may become manifest when the heart beats faster. Experimentally, under favourable conditions, e.g. when the heart is poisoned by certain drugs including digitalis, when it is

made to beat very fast or when its blood supply is curtailed short periods of alternation may follow a premature ectopic beat (Mackenzie 1907-8) or a dropped beat (Hering 1908) Sphygmograms show that pulsus alternans may begin abruptly either with an unusually large beat or with a small beat (Lewis 1925) and that the sum of 1 large and small beat equals the sum of two normal beats (Gaskell 188-) When the cardiac impulse appears to alternate in strength the peripheral pulse may behave concordantly or discordantly i.e. the larger pulse beat may be associated with the stronger or with the weaker cardiac impulse respectively (Hering 1908)

No thoroughly satisfactory hypothesis has been evolved to explain pulsus alternans It is generally believed that fewer muscle fibres contract with the weaker beats than with the stronger owing to the development of a state of partial refractoriness (Lewis 1925) fibres which do not contract with one beat recover in time for the next other fibres which contract with the first beat are still refractory and therefore unready for the second In other words there is a state of 2 : 1 partial ventricular response But if this were true all the beats should be weaker than normal the hypothesis does not explain the stronger beats Another suggestion is that pulsus alternans depends upon a disorder of ventricular relaxation for the ventricles hold more blood with the stronger beats and less with the weaker (Straub 1917)

Pulsus alternans should not be confused with electrical alternation (fig 506) nor with coupled beats due to premature systoles Electrical alternation is sometimes associated with pulsus alternans however as in the case illustrated

CHEYNE STOKES' BREATHING

Periodic breathing was described by Cheyne (1818) in what was probably a case of hypertensive heart failure with right hemiplegia For several days his breathing was irregular it would entirely cease for a quarter of a minute then it would become perceptible though very low then by degrees it became heaving and quick and then it would gradually cease again this revolution in the state of his breathing occupied about a minute Stokes (1854) connected the phenomenon with serious heart disease

Mechanism Cheyne Stokes breathing may be induced experimentally by hyperventilation especially in the presence of anoxia Over breathing washes out carbon dioxide and the ensuing apnoea is due to carbon dioxide lack During the apnoeic phase there is progressive anoxæmia until re-accumulation of carbon dioxide excites the respiratory centre and breathing is resumed During the dyspnoeic phase carbon dioxide is again washed out and the cycle repeats itself Anoxæmia and depression of the respiratory centre favour the production of Cheyne Stokes breathing The administration of carbon dioxide abolishes the giving of oxygen diminishes and modifies the periodicity The exact mechanism is unlikely to be understood until tissue chemistry is more advanced especially that relating to the

respiratory centre. The crescendo character of the dyspnoic phase may be partly due to time lag when respiration starts and carbon dioxide in the blood entering the lungs is blown off, blood which has already passed the pulmonary capillaries must have a higher carbon dioxide content than that which was required to galvanise the respiratory centre into action. This takes 5 to 10 seconds to reach the respiratory centre in normal subjects and an average of about 20 to 25 seconds in patients with left ventricular failure.

Clinical features Periodic breathing may result from a cerebral lesion, e.g. a head injury or a cerebral vascular accident, or from left ventricular failure usually in patients with hypertensive or ischaemic heart disease when sclerosis of cerebral vessels may be associated.

The cerebral type is characterised by a rise of blood pressure and pulse rate during the dyspnoic phase (Eyster 1906) in patients with left ventricular failure the central venous pressure and blood pressure rise during dyspnoea, the pulse rate and fore arm blood flow during apnoea (Sharpey Schafer 1948). Rhythmic variation in the size of the pupils may also be observed they dilate during dyspnoea and contract during apnoea.

Cheyne Stokes breathing is exaggerated by anything which further depresses the respiratory centre e.g. by morphine barbiturates, or natural sleep. It may cause insomnia by waking the patient at the height of the dyspnoic phase.

RIGHT VENTRICULAR FAILURE CONGESTIVE HEART FAILURE

When the right ventricle fails to discharge its contents adequately the pressure in the right auricle and the venae cavae rises the liver becomes enlarged and tender and dependent oedema usually develops.

ETIOLOGY

Right ventricular failure may result from massive pulmonary embolism subacute or chronic pulmonary heart disease pulmonary stenosis atrial septal defect or berri beri.

The term congestive heart failure is preferable when systemic congestion complicates mitral stenosis left ventricular failure rheumatic or other forms of carditis thyrotoxicosis or other hyperkinetic circulatory states (except those mentioned above) serious abnormalities of rhythm patent ductus arteriosus or other diseases affecting the heart as a whole. Failure in mitral stenosis is considered more fully on page 293 it is a mixture of pulmonary congestion due to a left sided lesion and right ventricular failure.

CLINICAL FEATURES

Elevation of the venous pressure By far the most important sign of right ventricular failure is a rise of systemic venous blood pressure. Its detection depends essentially upon clinical observation but direct measurement may

be employed as a check. Inspection of the cervical veins should be carried out with the subject horizontal or inclined at an angle of 30 45 60 or 90 degrees whichever position is most favourable. Venous pulsation may be distinguished from arterial in several ways. It is diffuse and undulant at least two waves being seen to each heart beat (except in certain special circumstances) whereas arterial pulsation is local abrupt and single. If the jugulars are compressed at the root of the neck venous pulsation ceases above this level whereas the carotids continue to beat. Abdominal compression increases the height and amplitude of jugular pulsation but has no effect on the carotids. The extent of cervical venous pulsation varies greatly according to the position of the patient whereas carotid pulsation scarcely alters. Finally the upper level of jugular pulsation may be seen to vary with respiration moving down with inspiration and up with expiration. The mean vertical height of this upper level above some arbitrary reference point such as the sternal angle should be measured in centimetres. The venous pressure may then be recorded in cm above the sternal angle with the patient at a known inclination. In normal subjects inclined at 45 degrees cervical venous pulsation is not seen at all but it may appear in the supraclavicular fossa at 30 degrees and may be 1 to 3 cm above the sternal angle in the horizontal position.



Fig 5 07- Photograph showing distension of the external jugular vein in case of congestive heart failure

Examination of the external jugular veins may also be helpful. If one of these vessels is constricted by the finger at the root of the neck a column of blood distends the upper part of the vein. On removing the finger the vein collapses. With the patient propped up at an angle of 45 degrees complete collapse of the vein at the root of the neck denotes no rise of venous pressure. On the other hand if the venous pressure is raised only the upper part of the vein collapses and it is easy to see the level at which pulsation ceases in the dilated lower part (fig 5 07). In assessing the venous pressure by observing the external jugulars it is important to make sure

they are pulsating for it is not unusual (nor abnormal) to find one or both of them dilated but not pulsating as a result of constriction at the point where they pierce the deep cervical fascia

The amplitude of venous pulsation is also worth noting for in pericardial effusion and chronic constrictive pericarditis it is diminished and in tricuspid incompetence it is enhanced

Occasionally the venous pressure is so high that pulsation which only occurs at the top of the column cannot be seen at all. If the veins are not obviously distended the raised venous pressure may then be overlooked or if recognised may be attributed to superior vena cava obstruction. Intracardiac catheterisation provides a valuable method of arriving at the truth. Superior vena cava obstruction is often only partial and the sudden fall in venous pressure as the catheter slips through the constriction is diagnostic.

Observation of cervical venous pulsation is still incomplete without noting its quality and its relationship to the arterial pulse. The *a* and *c'* waves of the jugular phlebogram may be difficult to detect with precision but it is usually possible to make out two waves and two troughs and it may be quite easy to time the main wave or trough.

A single abrupt collapsing type of venous pulsation in presystole denotes an exaggerated auricular *a* wave and is no evidence of failure. It alters little with change of posture, may be palpated (when it feels like a venous water hammer), and is sometimes transmitted to the liver as described by Mackenzie (1902). It is most conspicuous in tricuspid stenosis but may be associated with any condition which gives rise to powerful right auricular contraction (commonly cases with high right ventricular pressures and tall P waves). Cannon waves due to right auricular contraction against a closed tricuspid valve (as in heart block) look very similar but occur during ventricular systole.

In tricuspid incompetence the main venous wave is systolic and is usually powerful and prolonged. The normal depression following *c'* due to the sucking action caused by descent of the atrio ventricular septum is replaced by a wave of high pressure transmitted directly from the right ventricle thus the *c* and *v* waves become more or less fused. When there is auricular fibrillation the single forceful systolic venous wave so produced may be mistaken for an arterial pulse. Such a wave also is no direct evidence of congestive failure.

The chief venous wave in heart failure appears to be late diastolic. Strictly speaking it may not be a venous pulse wave at all but merely the steady rise of venous pressure that follows the momentary drop resulting from opening of the tricuspid valve. It is seen best in cases of auricular fibrillation when it merges into the *c* wave at the onset of systole and is followed by an abrupt systolic collapse due to descent of the base.

Clinical analysis of the venous pulse may not be easy but there can be no question that five minutes spent observing the movements of the neck veins may be as informative as auscultation.

Although elevation of the venous pressure at rest as described in previous paragraphs usually denotes congestive heart failure and is a constant sign of such there are certain circumstances in which it must be interpreted with caution. Slight elevation for example may occur when the intra-abdominal tension is high as in pregnancy, ascites and intestinal distension or when the intrathoracic pressure is raised from pleural effusion or pneumothorax. More important however are certain conditions (e.g. anaemia) in which elevation of the venous filling pressure is physiological as described on page 155. The venous pressure is also raised in hydremic states from any cause e.g. in acute nephritis and experimental water retention. Finally it may be very high in chronic constrictive pericarditis and in pericardial effusion.

When the venous pressure is within normal limits at rest it may yet rise unduly on slight exertion and may take several minutes to regain its resting level. This is a manifestation of limited cardiac reserve. The jugular venous pressure normally falls on exertion because increased ventilation lowers the mean intrathoracic pressure; the true filling pressure tends to rise.

Enlargement and tenderness of the liver. Hepatic distension may cause spontaneous pain in the right hypochondrium especially when it is rapid as in failure from paroxysmal tachycardia. Sometimes the pain is related to effort.

Palpation of the liver should be preceded by inspection and percussion. Epigastric fullness and dullness to percussion are characteristic of hepatic engorgement; on the other hand epigastric flattening or concavity with resonance to percussion are incompatible with it. Percussion of the right hypochondrium during the different phases of respiration often reveals the size of the liver with as much precision as palpation. The latter is best carried out with the left hand, the physician standing to the patient's left. It may be helpful to place the right hand high up under the right lower ribs and to exert forward pressure in order to push the liver towards the anterior abdominal wall. If the organ is distended its edge can be felt with the forefinger of the left hand as it moves downwards during inspiration. Pressure over an engorged liver is painful and causes immediate swelling of the cervical veins. Hepatic pulsation may be felt in cases of tricuspid incompetence expanding coinciding with the c wave of

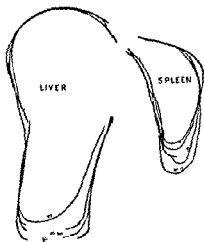


Fig. 508.—Tracings of serial skiagrams of liver and spleen opacified by means of thorotast demonstrating the rapid shrinkage of the liver and spleen which occurs when 1.5 mg. of digoxin is administered to a case of congestive failure.

the phlebogram. If there is ascites, an enlarged liver may be recognised by dipping—a repeated sudden pressure of the hand over the region of the liver when a sensation like that of a patella tap or like that of ballotting a foetus in utero may be appreciated. The liver shrinks as engorgement is relieved (fig 5 08) and this may be demonstrated within half an hour of giving 1.5 mg. of digoxin intravenously (Wood 1940). After repeated attacks of failure or after years of persistent distension, cirrhotic changes



Fig 5 09—Dependent oedema in congestive heart failure

may occur but they are unimportant and rarely interfere with hepatic function or with portal drainage.

Although distension and tenderness of the liver are useful signs of right ventricular failure, they are of secondary importance to elevation of the venous pressure upon which they depend. Bad diagnostic errors have been made when heart failure has been diagnosed on the combination of enlargement of the liver and dependent dropsy without a rise in venous pressure. In such cases carcinoma of the stomach or cirrhosis of the liver should receive first consideration.

Oedema. Of the three classic signs of congestive heart failure, oedema is the least reliable. It may be absent when the venous pres-

sure is high and gross when it is not so high. It is frequently absent in acute cases, especially in children. Cardiac oedema is essentially dependent (fig 5 09) but is occasionally observed in the face and is not infrequent in the arms. It is of course accompanied or preceded by oliguria and by a gain in body weight; in fact as much as six litres of fluid may collect in the tissue spaces before pitting oedema is necessarily demonstrable.

Physiologically it is thought that water, electrolytes and certain other small molecules such as sugar and urea leave the blood stream at the arterial end of the capillaries and re-enter at the venous end; the forces at work including the hydrostatic and osmotic pressures within and without the vessels and the permeability of the vascular endothelium. At the arterial end of the capillary the hydrostatic pressure exceeds the osmotic; at the venous end it is the other way about. The normal state of fluid balance may be upset in favour of the tissues by raising the hydrostatic

pressure within the capillaries or reducing it without by increasing the osmotic pressure within the capillaries or raising it without or by increasing the permeability of the vascular endothelium.

Increased hydrostatic pressure at the venous end of the capillaries is the cause of œdema in venous thrombosis, cirrhosis of the liver with tense ascites and in partial or complete obstruction of the superior vena cava. Low extra capillary pressure may determine the site of œdema but it does not cause it. Lax tissue occurs naturally in certain situations, e.g. in the infraorbital region and may be demonstrated subcutaneously following considerable loss of weight or when the skin has been stretched by previous dropsy. Reduction of capillary osmotic pressure is due mainly to reduction of plasma albumin. Œdema usually develops when the total blood proteins fall below 5 G per cent. Nephrosis, protein starvation, severe chronic anaemia and gross protein loss in pleural or peritoneal exudates may provide examples of such œdema. The chief effect of increasing the permeability of the capillaries is to allow more albumin to escape into the tissue spaces (a certain amount escapes normally and re-enters the blood stream via the lymphatics) and so to increase the osmotic pressure of the tissue fluid. Œdema with a high protein content (3 to 4 G per cent) results. Such œdema may be associated with burns, trench feet, insect bites and allergy (e.g. Quincke's disease). Lymphatic œdema has a similar high protein content.

The mechanism of the two most important forms of œdema, cardiac and nephritic, is not yet fully understood. In both, as a rule, the protein content of fluid samples is low (less than 1 G per cent), the venous pressure is raised and the blood volume is increased (Warren and Stead, 1944) but there are exceptions. Thus in chronic anaemia with congestive heart failure the blood volume is much diminished (Sharpey-Schafer, 1944). Increased capillary permeability is excluded by the low protein content of the œdema fluid; moreover, the theory that anoxia might be the cause of such capillary dysfunction is unlikely, in that cardiac œdema may be associated with a high cardiac output and normal arterial oxygen saturation, as in arterio-venous aneurysm. Elevation of the hydrostatic pressure at the venous end of the capillaries must play a part, but not necessarily a major part. In partial superior vena cava obstruction, for example, œdema does not occur until the venous pressure is very much higher than it is in heart failure. Reduction of renal blood flow to about 25 per cent of normal in most cases of congestive failure has been demonstrated (Merrill, 1946) and there is a considerable degree of sodium retention, according to Merrill and Cargill (1948), œdema occurs when the filtration rate falls below 70 to 80 ml./litre, tubular reabsorption being almost complete. Certainly artificial sodium retention, contrived by means of a high salt intake and desoxycorticosterone, may cause œdema, and cardiac dropsy is best relieved by means of a low sodium diet or sodium diuresis.

OTHER MANIFESTATIONS OF CONGESTIVE HEART FAILURE

It cannot be stressed too strongly that the diagnosis of congestive heart failure rests chiefly upon its peripheral effects and least upon central cardiac findings. In addition to the fundamental signs of failure already mentioned there are a number of other features which may be helpful in doubtful cases or which should be understood in order that their presence may not cause confusion. They include albuminuria and cylinduria, hydrothorax and ascites, cerebral symptoms, cardiac cachexia, venous thrombosis, jaundice, polychromasia, slowing of the erythrocyte sedimentation rate and certain radiographic appearances.

Urinary findings. Oliguria of course is associated with œdema. The urine which is rich in colour and of high specific gravity often contains albumin and hyaline casts and sometimes a few red cells.

Hydrothorax may occur from left or right ventricular failure and though usually bilateral tends to be left sided with the former and right sided with the latter (Bedford and Lovibond 1941). It should be remembered that the visceral pleura is drained by a venous plexus which is composed of both bronchial and pulmonary venous radicles. In typical instances the fluid is a transudate with a specific gravity ranging between 1.015 and 1.020, protein is often between two and three per cent and there may be moderate numbers of leucocytes and red cells. Unsuspected pulmonary infarction may further complicate the picture increasing the specific gravity, the protein content, the leucocyte count and especially the number of red cells, the overlying pleurisy giving rise to an exudate. If the fluid is frankly hæmorrhagic associated pulmonary infarction may be diagnosed with confidence.

Ascites is less common than hydrothorax and usually implies long standing failure. It is a special feature of tricuspid lesions and of chronic constrictive pericarditis.

Hydropericardium is rare and is usually of little significance. Cardiac compression does not occur, the electrocardiogram is uninfluenced and there are no symptoms. It is only important in that it alters the size and shape of the heart shadow and so may confuse radiographic observations.

Cerebral symptoms. Difficulty in concentration, impairment of memory, mental confusion, change of character and manic depressive, paranoid or other psychotic states are by no means rare accompaniments of heart failure. They are probably attributable to diminished cerebral blood flow, small cerebral thromboses or anoxæmia and are encountered particularly in hypertensive or ischæmic heart failure when cerebral arteriosclerosis may be partly responsible and in severe anoxic pulmonary heart disease especially when complicated by broncho-pneumonia.

Cardiac cachexia. Patients with chronic heart failure usually lose flesh although loss of weight may be prevented by fluid retention, thus wasting may only be noticed after diuresis, sometimes it is so great as to warrant the

term cachexia. Elevation of the basal metabolic rate, anorexia, impairment of intestinal function and enforced muscular inactivity may be partly responsible.

Venous thromboses are common in congestive heart failure especially when the cardiac output is low. They are responsible for the frequency of pulmonary infarction.

Jaundice may occur as a complication of severe heart failure and may be mainly obstructive (McMichael and Sherlock 1945) or mainly hemolytic, the former depending perhaps upon the raised intra hepatic pressure, the latter upon the destruction of red cells in hæmorrhagic pulmonary infarcts. The serum bilirubin is often in the region of 2 mg per cent.

Immature red cells are a common feature of congestive heart failure and may be due to stimulation of the bone marrow by anoxia.

The *erythrocyte sedimentation rate* is often retarded by congestive failure (Wood 1936). Figures of 50 to 100 in one hour obtained by the Westergren

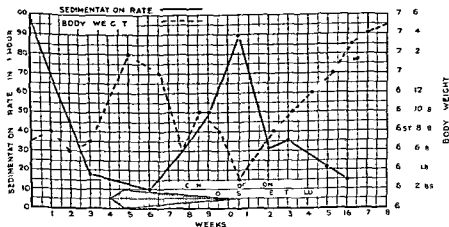


Fig 5 10—Fall in erythrocyte sedimentation rate resulting from the development of congestive failure in a case of active rheumatic carditis

method in cases of rheumatic carditis, myocardial infarction and syphilitic aortic incompetence may drop below 10 with the onset of failure and rise to their former level with recovery (fig 5 10).

Radiographic appearances The transverse diameter of the heart is increased by 1 to 2 cm during failure (figs 5 11a and b). In making such measurements care must be taken to exclude apparent enlargement due to raising of the diaphragm by an enlarged liver so that the heart takes up a more horizontal position. The superior vena cava throws a denser shadow than usual and the right auricle is more prominent. The lesser fissure on the right side may be clearly marked owing to pleural congestion or hydrothorax may be evident.

OTHER CONSIDERATIONS RELEVANT TO HEART FAILURE

Cyanosis Cyanosis is by no means a constant feature of heart failure and its presence in association with heart disease does not in itself indicate failure. It depends upon the presence of at least 5 G per cent of reduced hæmoglobin in the skin capillaries. Thus it cannot occur in severe anaemia but develops readily if there is polycythæmia. The intensity of the hue depends upon the calibre of the skin capillaries: if they are constricted the colour is paler; if dilated it is richer. Polycythæmia is the rule in the cyanotic forms of congenital heart disease and it occurs occasionally in pulmonary heart disease. Capillary dilatation may be seen in the face in many cases of mitral stenosis.

There are three principal causes of cyanosis in heart disease. The first is the right to left shunt seen in congenital heart disease when venous blood passes directly into the arterial circulation. Central cyanosis of this kind cannot be recognised clinically unless at least one quarter of the venous blood passes through the defect assuming a normal hæmoglobin value. With a hæmoglobin of 120 per cent only one fifth of the cardiac output need be shunted to produce cyanosis and with a hæmoglobin of 140 per cent only about one sixth. The second cause is inadequate oxygenation of blood passing through the lungs owing to failure of alveolar function and is chiefly encountered in acute pulmonary oedema and in pulmonary heart disease secondary to emphysema. Clinical recognition of such central cyanosis means that the arterial oxygen saturation is reduced to 80 per cent or less if the hæmoglobin value is normal. The third cause is the most common and is the sluggish peripheral blood flow in the skin due to compensatory vasoconstriction. It is seen especially in mitral stenosis, primary pulmonary hypertension and massive pulmonary embolism but it may occur in congestive heart failure from any cause provided the cardiac output is low.

Considerable clinical difficulty may be experienced in attempting to distinguish between central and peripheral cyanosis. If the skin is cold cyanosis of the face, ears and nail beds must be assumed to be due to peripheral vasoconstriction; if the skin is warm and especially if a water hammer pulse, digital throbbing and capillary pulsation can be demonstrated cyanosis is probably central. The colour of the conjunctiva, the inside of the lips or of the palate may be more informative for cyanosis in these warm situations is always central. Direct measurement of the arterial oxygen saturation is recommended in all cases of doubt if accurate information is desired.

The administration of oxygen is most valuable in cases of central cyanosis due to emphysema but it also increases the arterial oxygen saturation in cyanotic cases of congenital heart disease and may be used to tide such cases over some critical period.

Behaviour of the blood pressure The blood pressure might be expected to fall in congestive heart failure but in fact it may rise fall or remain stationary in the majority of cases it rises. There are only two conditions in which heart failure is characteristically associated with a sharp drop of blood pressure acute myocardial infarction and massive pulmonary embolism. In the former the drop is not related to failure (see page 389) in the latter it is more or less proportional to the reduction in cardiac output and hence to the size of the embolus. Conspicuous lowering of the blood pressure associated with heart failure in other diseases is commonly a terminal event. The vasoconstriction that maintains the blood pressure when the cardiac output falls may depend upon diminished blood flow through the kidney. It may be recognised clinically by cold extremities and peripheral cyanosis.

The heart rate Whilst some degree of tachycardia partly due to the Bainbridge reflex (page 110) is usual in heart failure there are extreme variations both with normal and abnormal rhythms. If the heart rate is plotted against the cardiac output a curve may be constructed which is more or less similar in shape to that related to the venous pressure. To some extent it is likely that tachycardia in heart failure represents another compensatory device whereby the cardiac output may be increased. This is well illustrated in chronic constrictive pericarditis when the venous pressure mechanism fails. Clinically however tachycardia is an unreliable guide to the presence or degree of failure as the pulse rate is influenced by so many other factors.

Character of the heart sounds Current terminology still includes such expressions as weak faint or distant heart sounds and tic tac or fetal rhythms which have been supposed to signify failure or threatened failure. With the exceptions of the reduction in the intensity of the heart sounds following coronary thrombosis pulmonary embolism and pericardial effusion weak faint or distant heart sounds are commonly due to obesity emphysema or well developed thoracic muscles. It is doubtful whether tic tac or fetal rhythm is in any way associated with central heart failure on the other hand it is heard in patients suffering from shock and may be associated with diminution of the blood volume. A weak first heart sound associated with a normal second sound is usually due to a P-R interval around 0.21 to 0.22 second the mitral cusps then having time to float into apposition before the ventricles contract (Levine 1948).

PROGNOSIS OF HEART FAILURE

When left ventricular failure develops in the natural course of hypertensive or aortic valve disease the prognosis is poor the patient seldom living more than eighteen months after the onset of orthopnoea or paroxysmal cardiac dyspnoea. Few die however before clinical signs of chronic systemic congestion become apparent. Moreover right ventricular failure often brings symptomatic relief as it reduces pulmonary congestion.

The prognosis may be less unfavourable when acute myocardial infarction is responsible because if the patient survives the acute phase he may make a good recovery and although the average life expectancy is still only 3 to 4 years the chances of much longer survival are not remote.

The outlook is entirely different when left ventricular failure complicates acute nephritis here complete recovery may be anticipated. The ultimate prognosis depends upon the subsequent course of the nephritis. Similar remarks apply to other forms of hypertension which are transient or which can be treated successfully.

The prognosis of right ventricular failure or congestive heart failure depends very much upon its cause. When associated with diseases that can be cured or improved such as thyrotoxicosis the outlook is excellent. On the other hand when it occurs in the natural course of chronic and incurable heart disease few patients survive more than a year. Between these extremes are cases of incurable heart disease in which failure is precipitated by some adverse factor which is either transient or which can be improved or cured. Undue physical work, pregnancy, infection, disturbances of rhythm, obesity and pulmonary embolism provide examples of such factors.

IRITANT

As the measures used in the treatment of left and right ventricular failure are practically the same they will be considered together. By failure is meant the final stage in which the heart is overloaded.

Rest in bed or in a comfortable armchair is essential and should be continued for a minimum period of three weeks. If signs of failure do not disappear within a few days of instituting adequate therapy the period of rest should be extended to six weeks. The patient should be nursed against a back rest at an angle of about 60 degrees whether orthopnoic or not for there is no easier way of lowering the right auricular pressure and so unloading the overburdened heart if the legs are lowered so much the better—hence the value of an armchair. *Meals* should be small in quantity and fluids limited to about two pints daily. If the *sodium intake* can be limited to 0.5 G daily however there is no need to restrict fluids. Correct treatment of heart failure usually serves as the best hypnotic but if insomnia is troublesome at first there should be no hesitation in using morphine.

Venesection deserves a better reputation. It has fallen out of favour because similar results may be obtained by means of certain drugs but it offers a quick and sure way of lowering the venous pressure and reducing hydræmia and should not be abandoned. Drugs which lower the venous pressure should be used in addition not as a substitute. About 600 to 750 ml. of blood may be withdrawn.

Digitalis is beneficial whether there is auricular fibrillation or normal rhythm and whether the pulse rate is fast or slow. It lowers the venous pressure (fig. 5 12) raises the blood pressure (fig. 5 13) slows the heart rate

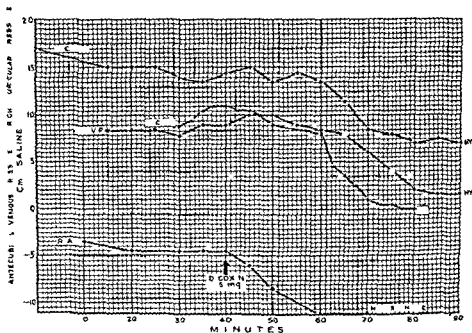


Fig 5 12—Typical effect of digitalis on the venous pressure or right auricular pressure in four cases of congestive heart failure

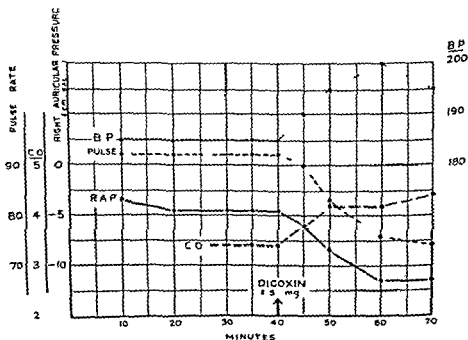


Fig 5 13—Typical effect of digitalis on the blood pressure, pulse rate, right auricular pressure and cardiac output in a case of hypertensive heart failure with normal rhythm

ACUTE RHEUMATIC CARDITIS CONGESTIVE HEART FAILURE

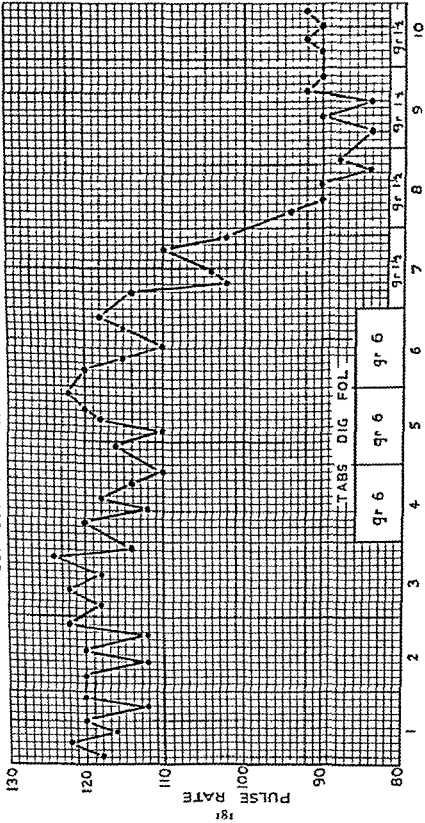


Fig. 5-14-67 showing the progression of pulse rate in a case of acute rheumatic carditis and congestive heart failure.

DIGITALIS IN LEFT VENTRICULAR FAILURE

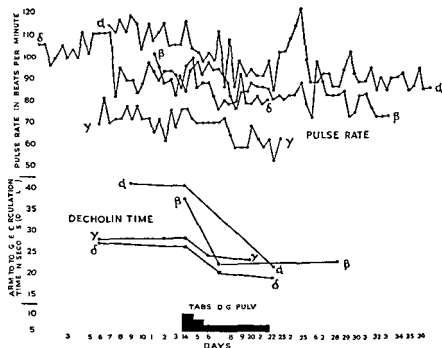


Fig 5 15—The action of digitalis on the arm to tongue circulation time and on the pulse rate in four cases of left ventricular failure with normal rhythm

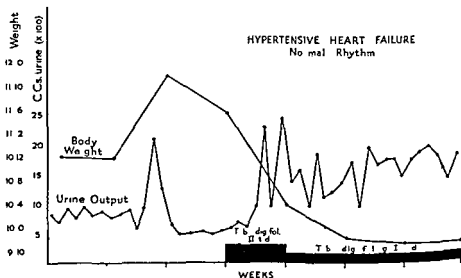


Fig 5 16—Chart showing considerable diuresis resulting from the administration of digitalis to a case of hypertensive heart failure with normal rhythm

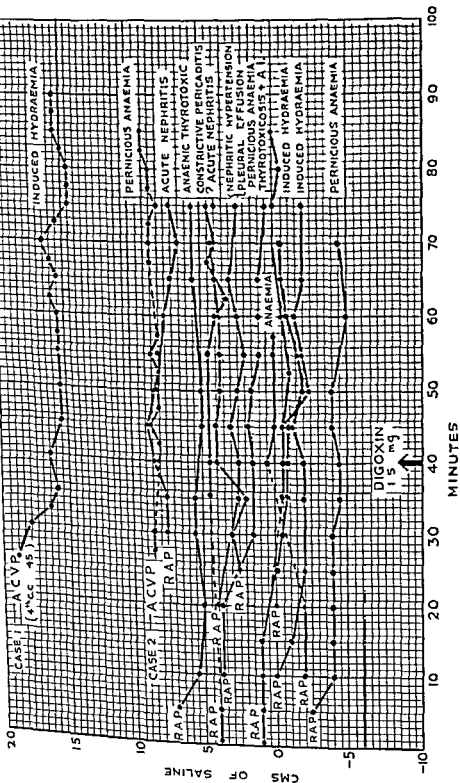


Fig 5 17--Chart illustrating the failure of digitalis to lower the right auricular pressure in twelve cases in which it was raised from causes other than congestive failure

(fig 5 14) relieves hepatic distension (fig 5 08) increases the vital capacity, shortens the pulmonary circulation time (fig 5 15) increases the cardiac output (fig 5 13) and encourages diuresis (fig 5 16) Its good effects have been recently attributed to a direct venous pressure lowering action (McMichael and Sharpey Schafer 1944) but this is doubtful for digitalis does not lower the venous pressure when the latter is elevated from causes other than congestive heart failure (fig 5 17) (Wood and Paulett 1949) The original belief that digitalis improves the function of the heart by virtue of its direct action on the myocardium is probably correct In normal controls increase of myocardial tone may make the heart smaller and may reduce its output (Stewart *et al* 1938)

For routine purposes the dose of digitalis should be 3 grains (0.2 G) of the powdered leaf *t d s* on the first day 2 grains (0.13 G) *t d s* on the second and 1 grain (65 mg) *t d s* thereafter until demonstrable improvement or evidence of intoxication occurs when it may be reduced to 1 grain (65 mg) *b i d* Other methods of administering digitalis are described on pages 144 and 147

Strophanthin may be preferred when a quick action is desired especially if a cumulative effect is not wanted A single dose of Ouabain 1.0 mg intravenously may raise the cardiac output in cases of heart failure without affecting the venous pressure (McMichael 1948) and so presumably acts directly on the heart Like intravenous digoxin it also has a conspicuous pressor effect and slows the pulse rate Strophanthin is probably the drug of choice in collapsed cases of pulmonary heart failure

Mercurial diuretics provide the best means of reducing œdema More over proportional and parallel to the diuresis and to the relief of hydremia the venous pressure falls For this reason they also prevent paroxysmal cardiac dyspnoea (fig 5 18) They act by encouraging sodium excretion

Preparations on the market include mersalyl mercuraphylline salyrgan neptal and novurt They are all based on the original but far more toxic substance novarsurol and contain about 40 per cent mercury Ampoules for injection contain 10 per cent of the drug and 5 per cent of theophylline oral tablets 0.08 G of the mercurial diuretic and 0.04 G of theophylline and suppositories 0.4 G of mersalyl and 0.2 G of theophylline

Mersalyl should be given in doses of 2 ml intramuscularly twice weekly accompanied by 30 grains (2 G) of ammonium chloride orally once on the preceding evening and repeated three times on the day of injection Mercurial diuretics may be given orally in doses of two tablets *t d s* for two days with a rest of three or four days between courses but they are less effective by this route and may cause considerable gastro intestinal disturbance Rectal suppositories one per week may also be used but may provoke severe burning pain

Toxic reactions are rare but sudden death has been reported after intravenous injections Toxic nephrosis characterised by tubular degeneration and calcification is encountered occasionally usually after prolonged

administration. The drug should not be stopped owing to a poor initial response for the result of the second or third dose coinciding perhaps with the beneficial effect of rest and digitalis may exceed expectations. The only contra-indication is acute nephritis.

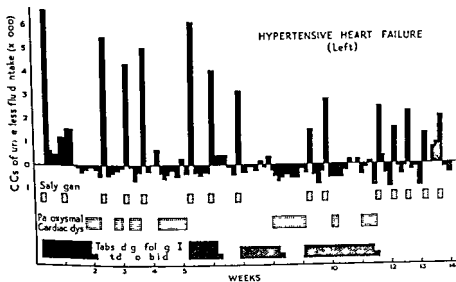


Fig. 18—Chart illustrating the beneficial effect of mercurial diuretics in preventing paroxysmal cardiac dyspnoea. Digitalis was less effective.

Other diuretics which may be employed as adjuvants include the xanthine derivatives theobromine, theophylline and caffeine and urea in massive doses e.g. 30 G t.d.s. The most powerful of this group is theobromine which is given in doses of 5 to 10 grains (0.32 to 0.65 G) or combined with sodium salicylate as diuretin in doses of 10 to 20 grains (0.65 to 1.3 G). The latter being far more soluble is preferable.

A *low sodium diet* has proved a most effective way of relieving obstinate oedema (Schroeder 1941) and preventing paroxysmal cardiac dyspnoea. The object is to reduce the sodium intake to the order of 0.5 G daily so that it is impossible for the tissues to hold much fluid. The blood volume is thus reduced and the venous pressure lowered. The following diet has been constructed from tables giving the composition of numerous foods compiled by McCance and Widdowson (1946). The first figure after each substance gives the amount of sodium in mgs. per 100 G of foodstuff. The second figure gives the approximate calorific value of the food per mg of sodium content. For the first 48 hours it is a good plan to give nothing but fruit in any form, fruit juice, drinks, sugar, rice and diluted milk. Mercurial diuretics should not, as a rule, be given in conjunction with this diet, the combination causing a too profound degree of sodium and chloride depletion, uræmia which may prove fatal may then develop.

LOW SODIUM DIET

CEREALS

<i>Permitted</i>			<i>Doubtful</i>			<i>Forbidden</i>		
Arrowroot	48	72	Current bread	164	2	Bread	393	07
Barley	08	150	Sweet biscuits	216	3	Biscuits	400	08
Cornflour	52	7	Rusks	200	2	Cornflakes	1050	03
Flour	25	170				Grapenuts	658	05
Macaroni	79	15				Post Toasties	810	05
Oatmeal	33	11				Ryvita	615	05
Rice	22	60				Vita wheat	615	05
Sago	34	100						
Semolina	12	30						
Shredded								
Wheat	16	22						
Tapioca	4	86						

NOTE

Biscuits Water biscuits and cream crackers contain the most sodium. Oatmeal biscuits made without salt and with lard instead of margarine are recommended.

Breakfast cereals Oatmeal porridge should be made without salt and with equal parts of milk and water. Shredded wheat with diluted milk and plenty of sugar is recommended.

Milk puddings Milk should be diluted with equal parts of water. margarine must not be used.

Flour sauces Make without salt and with equal parts of milk and vegetable water. Use dripping instead of margarine.

DAIRY PRODUCE AND FATS

<i>Permitted</i>			<i>Doubtful</i>			<i>Forbidden</i>		
Butter (fresh)	223	35	Milk (fresh)	50	12	Cheese	600	05
Cream cheese			Milk (sweet condensed)	143	2	Egg white	192	02
(home made)	110	8				Margarine	318	05
Cream	31	13						
Egg yolk	50	7						
Olive oil	01	9290						
Lard	2	450						
Dripping	5	200						
Suet	25	44						

NOTE

Butter may be kneaded in water to reduce its salt content.

Home made cream cheese must be made without salt.

Eggs are best fried or poached because a certain amount of sodium chloride is then lost in the cooking.

Dilute milk 2:1

Use olive oil, dripping, lard or suet in cooking instead of butter or margarine whenever possible.

MEAT, POULTRY AND GAME

<i>Permitted</i>			<i>Doubtful</i>			<i>Forbidden</i>		
Roast beef	62	6	Chicken	80	2	Bacon	1200	03
Grilled steak	67	5	Duck	195	15	Beef		
Stewed steak	38	55	Goose	145	2	(silver side)	1470	02
Wate (roast or stewed)	45	45	Guinea fowl	130	15	Brains	150	07
			Heart	153	15	Ham	1500	03

<i>Permitted</i>		<i>Doubtful</i>		<i>Forbidden</i>	
Mutton chop (grilled or fried)	90	Liver	100	Kidney	250
Mutton leg etc (roast boiled or stewed)	68	Partridge	25	Meat paste	940
Pork roast	66	Pheasant	100	Smoked pork	1800
Pork chops	60	Pigeon	2	Sausage	1000
Rabbit	32	Turkey	130	Tongue	180
Sweetbread	69	Tripe	172	(preserved)	180
Tongue (fresh)	79	Veal	100		
Topside (beef)	50	Venison	86		

NOTE

All salted and preserved meats are forbidden

Roasts are best as there are more calories per mg of sodium content

Meat extracts like Marmite Bovril and Oxo are forbidden

The simple meats - beef mutton lamb pork hare and rabbit - are the best

Next comes game Of offal sweetbread fresh tongue and liver are best

FISH

Steamed fish is mostly about 100/1

Fried fish slightly better (125/15)

The fish of choice are bass conger dabs silver eels herring red mullet pollan
fresh salmon fried smelts sprats and whitebait

Preserved smoked salted or tinned fish are prohibited (e.g. tinned salmon is
538/025)

Fish not advised include crab haddock flat fish lobster mussels oysters
(505/01) sea trout whiting and winkles

Fish paste is prohibited

Fish cakes made without salt and deep fried in olive oil are recommended

FRUIT

<i>Permitted</i>		<i>Permitted</i>	
Apples	2	Greengages	14
Apricots	1	Oranges	29
Bananas	12	Peaches	27
Blackberries	37	Pears	23
Cherries	28	Pineapple	17
Currants	27	Plums	17
Dates	47	Quinces	32
Figs	16	Raspberries	25
Gooseberry	12	Rhubarb	15
Grapes	16	Strawberries	17
Grape fruit	14		

NOTE

These are average samples of fresh fruits

The only doubtful fruits are melon (195/1) and passion fruit (30/1)

Stewed fruit is best because of its higher calorific value e.g. stewed apples
(01/170)

Tinned fruits in syrup are also good

Dried fruits are less beneficial e.g. tinned apricots (09/62) dried apricots
(56/3)

Preserved olives (2250/005) are forbidden

NUTS

Almonds	5 8	100
Brazils	1 5	430
Chestnuts	10 9	16
Hazelnuts	1 4	80
Walnuts	2 7	353

NOTE

These are average examples of fresh nuts

Obviously salted almonds and peanuts are forbidden

VEGETABLES

<i>Permitted</i>			<i>Doubtful</i>			<i>Forbidden</i>		
Artichokes (root)	6	7 3	Artichokes globe	6 4	1 1	Beetroot	64	0 7
Asparagus	0 9	10	Broad beans	19 6	2	Carrots	50	0 3
Butter beans	16 2	6	Cabbage	10	0 6	Celery	137	0 07
French beans	3 4	2	Cauliflower	11	1	Radishes	59	0 25
Harcot beans	15	6	Cucumber	13	0 7	Spinach	123	0 2
Runner beans	3 3	2	Mustard and cress	10	0 5			
Sprouts	7 7	2	New potatoes	40 5	2			
Leeks	6 4	4	Swedes	14 4	1			
Lentils	9 4	10	Turnips	28	0 5			
Lettuce	3 1	3 5						
Marrow	1 2	6						
Mushrooms fried	11	20						
Onions boiled	6 6	2						
raw	10 2	2						
fried	20	18						
Parsnips	4 1	14						
dried	12 6	8						
Potatoes boiled	3 4	25						
roast	8 6	14						
Tomatoes	8	5						
Tomatoes fried	3 3	21						

NOTE

Vegetables must be cooked free from salt. They must not be mashed with margarine or salted butter.

Tinned or otherwise preserved vegetables e.g. tinned peas (260 0 3) are banned.

SWEETS

Sugar (0 4 984) adds a low sodium high caloric value to most sweets.

Plain chocolate (18 6 29) is better than milk chocolate (93 4 6).

Honey (10 7 26) and jam (13 9 16) are recommended.

Golden syrup (270 1), chutney (130 1) and mincemeat (200 0 5) are prohibited.

Toffee (115 3 5) and black treacle (96 2 5) should be avoided.

BEVERAGES

<i>Permitted</i>			<i>Prohibited</i>		
Coffee	0 3	15	Bournvita	360	1
Lemonade	0 5	100	Bovril	5 280	0 02
Tea	0 4	2	Cocoa	650	0 7
Beer	15	3	Horlicks	690	0 6
Wine			Marmite	6 130	0 01
Spirits			Ovaltine	249	1 5
			Oxo Cubes	10 600	0 02
			Virol	374	1

CONDIMENTS

	<i>Permitted</i>			<i>Prohibited</i>	
Ginger	34	7 5	Curry	450	0 5
Mustard	5	90	Salt	38 500	0
Pepper	7	45			
Vinegar	20	0 2			

CAKES AND PASTRIES

Most of these work out at 150 3 approx

Doughnuts (60'6) oatmeal biscuits made without salt and with lard instead of margarine shortbread (86'6) sponge cake (79 4) apple charlotte with suet on top instead of margarine apple dumpling (39 5) apple pudding (48 5) jelly (8 9 5) pancakes (88'4) and cereal puddings made with diluted milk and no margarine are permitted

Foods made with soda bicarbonate are not allowed (e.g. dumpling)

Yorkshire pudding made without salt is permissible

Currant cake ginger cake and Swiss roll should be avoided

GENERAL RULES

No free salt or ordinary salt substitutes no salt in cooking sodium free salt substitutes usually made with potassium such as neo seleron are permitted

No foods made with baking powder

No medicines containing sodium

No preserved salted smoked or tinned foods (except dried and tinned fruit)

Dilute milk with equal parts of water

Use dripping lard olive oil or suet instead of butter or margarine wherever possible

Supply calories chiefly with selected cereals cream fat fresh meat potatoes sugar sweets fruit and nuts

Avoid bread bread substitutes certain cereals margarine salted butter cheese bacon sausages meat extracts shell fish fish paste certain vegetables and milk beverages

Acupuncture When œdema is gross and fails to respond to the measures previously outlined it may be necessary to resort to acupuncture. A triangular cutting needle is used and about a dozen punctures are made in each leg the patient is then seated in a chair with his legs in a tub. To facilitate drainage the legs may be swabbed down with warm citrate solution from time to time. Due antiseptic precautions must be maintained. Fluid may continue to exude for twenty four to forty eight hours and it is not uncommon for the total quantity to be measured in gallons. Southey's tubes constitute a cleaner way of removing fluid on the same principle. Several large bore needles are inserted into the subcutaneous tissue of the thighs or calves and fluid is allowed to drain away through attached rubber tubes into a container.

Attacks of paroxysmal cardiac dyspnoea or of acute pulmonary œdema are treated by methods designed to lower the venous filling pressure as quickly as possible and so to reduce the output of the right ventricle. The

sitting position will usually have been adopted already by the patient. Morphine $\frac{1}{4}$ to $\frac{1}{2}$ of a grain (16 to 21 mg) intramuscularly or $\frac{1}{4}$ of a grain (11 mg) intravenously depresses the excited respiratory reflexes and soothes the patient. Venous tourniquets may be applied round the thighs to trap blood in the legs or venesection may be preferred. Theophylline ethylenediamine (aminophylline) 0.24 to 0.48 G intravenously lowers the venous pressure immediately, relieves bronchial spasm and may have a direct stimulating action on the heart (fig. 502). Tetraethylammonium bromide 200 to 300 mg intravenously is a useful agent for lowering venous pressure and may relieve attacks quickly (Hayward 1948).

Digoxin and strophanthin are probably best avoided in view of their pressor actions; indeed paroxysmal cardiac dyspnoea may occasionally be initiated by intravenous digoxin.

Oxygen is of little value in paroxysmal cardiac dyspnoea for the arterial oxygen saturation is normal but may be given with advantage in acute pulmonary oedema. Nikethamide is contraindicated for the aim is to depress respiration not to stimulate it. Adrenaline is dangerous in ischaemic cases because it may provoke angina pectoris, paroxysmal ventricular tachycardia or ventricular fibrillation but it may be given in small doses subcutaneously to relieve bronchial spasm in hypertensive cases. Atropine should be avoided for it has no therapeutic value and causes unnecessary tachycardia.

Cheyne Stokes breathing may be abolished by carbon dioxide and relieved by oxygen as already stated but is best treated by an intravenous injection of 0.24 to 0.48 G of theophylline ethylenediamine. The effect lasts for six to eight hours and thus ensures a good night's rest. According to Marais and McMichael (1937) it is the ethylenediamine radical which is responsible but other workers (e.g. Nathanson and Fitzgibbon 1939) have reported equally good results with theophylline alone or with other theophylline salts. These conflicting findings are not necessarily contradictory for both may be effective: the ethylenediamine radical by direct action on the respiratory centre, theophylline by improving the state of the circulation.

Theophylline may also be given by mouth in doses of 0.2 to 0.3 G four hourly not only to prevent paroxysmal cardiac dyspnoea and Cheyne Stokes breathing but also to lower the venous pressure in congestive heart failure. Larger doses (0.4 to 0.5 G) may be tried but usually have to be abandoned owing to dyspepsia.

Dramatic results may follow treatment directed against the cause of the underlying heart disease. This applies particularly to cases of thyrotoxicosis, anaemia, beri beri, arterio-venous aneurysm and anoxic pulmonary heart disease.

If in spite of all these measures heart failure continues an attempt may be made to reduce the oxygen requirement and therefore the work of the heart by means of *thiourea* or total ablation of the thyroid gland. The former

is preferable because the treatment can be abandoned if unsuccessful. Relatively large doses are necessary, usually 0.6 to 0.9 G daily. Propylthiouracil is advised for it is less toxic than other preparations. It must be admitted, however, that results are far from satisfactory. Radioactive iodine offers another means of inducing artificial myxoedema (Blumgart *et al.* 1950).

REFERENCES

- Bedford D E and Lovibond J L (1941) Hydrothorax in heart failure *Brit Heart J* 3 93
- Blumgart H L, Freedberg A S and Kurland G S (1950) Hypothyroidism produced by radioactive iodine *Circulation* 1 1105
- and Weiss S (1927) Studies on the velocity of blood flow. The velocity of blood flow in the systemic and pulmonary circulations in health and disease *J clin Invest* 4 15 149 173 199 389 399 (1928) 5 343 379
- Cheyne J (1818) A case of apoplexy in which the fleshy part of the heart was converted into fat *Dublin Hosp Rep* 2 216
- Eyster J A E (1906) Clinical and experimental observations on Cheyne-Stokes respiration *Johns Hopk Hos Bull* 8 232
- Fishberg A M (1939) Hypertension and nephritis 4th ed. London
- Gallavardin L (1913) Pseudo Deboulement du Deuxieme Bruit du Coeur Simulant de Doublement mitral *Lyon Med* 121 409
- Gaskell W H (1882) On the rhythm of the heart of the frog and on the nature of the action of the vagus nerve *Phil Trans Roy Soc* 173 993
- Gibson A G (1907) The significance of a hitherto undescribed wave in the jugular pulse *Lancet* ii 1380
- Gross H and Spark C (1937) Coronary and extra coronary factors in hypertensive heart failure *Amer Heart J* 14 160
- Harrison T R (1935) Failure of the circulation. Baltimore
- Hayward G W (1948) Tetraethyl ammonium bromide in hypertension and hypertensive heart failure *Lancet* i 18
- Hering H E (1908) Das Wesen des Herzalternans *Munch med Wschr* 55 ii 1417
- Heyer H E, Holman J and Shires G T (1948) The diminished efficiency and altered dynamics of respiration in experimental pulmonary congestion *Amer Heart J* 35 463
- Hope J (1832) A treatise on the diseases of the heart. London
- Levine S A (1948) Auscultation of the heart. St Cyres lecture. National Heart Hospital, London
- Lewis T (1925) The mechanism and graphic registration of the heart beat. London
- (1933) Diseases of the heart. 1st ed. London
- McCance R A and Widdowson E M (1946) The chemical composition of foods. London
- Mackenzie J (1902) The study of the pulse. London
- (1907-8) The extrasystole: a contribution to the functional pathology of the primitive cardiac tissue *Quart J Med* 1 481
- (1913) Diseases of the heart. 3rd ed. London
- McMichael J (1939) Hyperpnoea in heart failure *Clin Sci* 4 19
- (1947) Circulatory failure studied by means of venous catheterisation *Advances in Internal Medicine* 2 64
- (1948) Pharmacology of the failing human heart *Brit med J* 1 927
- and Sharpey-Schafer E P (1944) The action of intravenous digoxin in man *Quart J Med* 37 1-3

- (1944) Cardiac output in man by a direct Fick method *Brit Heart J* 6 33
- and Sherlock S P V (1945) Jaundice in heart failure *Ibid* 14 222
- Marais O A S and McMichael J (1937) Theophylline ethylene diamine in Cheyne Stokes respiration *Lancet* 437
- Merrill A J (1946) Oedema and decreased renal blood flow in patients with chronic congestive heart failure Evidence of forward failure as primary cause of oedema *J clin Invest* 25 389
- and Cargill W H (1948) The effect of exercise on the renal plasma flow and filtration rate of normal and cardiac subjects *Ibid* 27 272
- Müller W S (1937) *The Lung* London
- Nathanson M H and Fitzgibbon J P (1939) Pharmacology of Cheyne Stokes Respiration *Amer Heart J* 17 691
- Ohm R (1913) Venenpuls und Herztone *Dtsch Med Wsch* 39 1493
- Potain P C (1876) Concerning the cardiac rhythm called gallop rhythm *Bull et mem soc med d Hop de Paris* 12 137
- Scadding J G and Wood P H (1939) Systolic clicks due to left sided pneumothorax *Lancet* ii 1258
- Schroeder H A (1941) Studies on congestive heart failure *Am Heart J* 22 141
- Sharpey Schafer E P (1944) Cardiac output in severe anaemia *Clin Sc* 5 125
- (1948) Personal communication
- Starling E H (1918) The Linnæus lecture on the law of the heart London
- Stewart H J Deitrick J E Crane N F and Wheeler C F (1938) Action of digitalis in uncomplicated heart disease *Arch intern Med* 62 569
- Stokes W (1854) On Fatty Degeneration of the Heart Chap 3 of *The diseases of the heart and Aorta* 1st ed Dublin
- Straub H (1917) Dynamik des Herzalternans *Deutsch Archiv klin Med* 123 403
- Traube L (1872) Ein Fall von Pulses Bigeminus nebst Bemerkungen über die Leberschwellungen bei Klappenfehlern und über acute Leberatrophie *Berl klin Wochenschr* 9 185
- Warren J V and Stead E A Jr (1944) Fluid dynamics in chronic congestive heart failure interpretation of mechanisms producing oedema increased plasma volume and elevated venous pressure in certain patients with prolonged congestive failure *Arch intern Med* 73 138
- Wood P H (1936) The erythrocyte sedimentation rate in diseases of the heart *Quart J Med* 5 1
- (1936) Right and left ventricular failure A study of circulation time and venous blood pressure *Lancet* ii 15
- (1940) The action of digitalis in heart failure with normal rhythm *Brit Heart J* 2 132
- and Laulett J (1949) The action of digitalis on the venous pressure *Ibid* 11 83

CHAPTER VI

SYNCOPE

THERE are many causes of transient loss of consciousness and a complete list would include the causes of epilepsy coma concussion and asphyxia but syncope has come to mean transient loss of consciousness of sudden onset due to inadequacy of the cerebral blood flow As so defined syncope may be divided into cardiac vasomotor or vaso-vagal cerebral and anoxic forms

CARDIAC SYNCOPE

Cardiac syncope occurs when the heart through some fault in itself or in its great vessels fails to maintain an adequate cerebral circulation These faults are listed for convenience as follows

- 1 Cardiac standstill – vagal inhibition
- 2 Ventricular asystole – Stokes Adams fit
- 3 Ventricular fibrillation
- 4 Ball valve thrombus or pedunculated myxoma
- 5 Aortic stenosis
- 6 Paroxysmal rhythm changes with extremely rapid ventricular rates
- 7 Massive pulmonary embolism
- 8 Cardiac compression from hæmopericardium

The immediate cause of such syncope is a sudden fall in cardiac output The practical mechanism whereby the heart fails to fulfil its task varies according to the lesion

In *cardiac standstill ventricular asystole ventricular fibrillation ball valve thrombus* and *pedunculated myxoma* loss of consciousness is abrupt and without warning The attack may occur at any time while the patient is walking standing sitting or lying At first the patient is grey or white flaccid pulseless and motionless The heart sounds are inaudible but respirations may continue In about 10 to 15 seconds anoxic twitchings begin and may develop into convulsions if the attack lasts long enough If recovery does not occur within two minutes death usually results Cardiac and ventricular asystole usually recover well within that time commonly within 5 to 10 seconds but ventricular fibrillation is usually though not necessarily fatal Ball valve thrombus and pedunculated myxoma are rare Return to consciousness is abrupt and complete and is followed by a vivid flush hyper-oxygenated blood being flung into a dilated vascular system (reactive hyperæmia)

Similar attacks of uncertain mechanism may occur in *aortic stenosis* As

a rule however syncope in aortic stenosis is vasomotor the valve lesion acting merely as a predisposing factor

Heart rates up to 200 per minute in *paroxysmal tachycardia* are usually well tolerated, but syncope may result if the rate is much faster. Speeds of over 300 per minute have been recorded

Massive pulmonary embolism may cause syncope when more than two thirds of the circulation is blocked. The onset is sudden but rarely so abrupt as in the group just mentioned. Moreover it may be preceded by pain or tightness in the chest. The duration of unconsciousness is longer being usually measured in minutes or even hours. Recovery is at first only partial extreme faintness persisting. During the attack the patient is limp grey sweating and breathless. The pulse is thready or imperceptible, the heart sounds faint or inaudible, the blood pressure low or unobtainable

Smaller pulmonary emboli insufficient seriously to embarrass the circulation occasionally cause reflex syncope (page 449). Such reactions may be prevented by means of atropine. Similar attacks may be encountered in cases of acute myocardial infarction. These should not be regarded as examples of cardiac syncope for the mechanism is vasomotor

Cardiac compression must be gross to reduce the cardiac output sufficiently to cause loss of consciousness. This condition may be fulfilled by hæmopericardium due to rupture of an aneurysm dissecting or saccular or to perforation of the heart from bullet or stab wounds or spontaneously through a myocardial infarct or ventricular aneurysm

Syncope associated with aortic incompetence is usually vasomotor in origin the lesion acting only as a predisposing factor for the peripheral resistance is already low

From this brief survey it will be seen that syncope associated with cardiac disorder is of two main types that in which there is an abrupt and gross fall in cardiac output (true cardiac syncope) and that in which the heart lesion predisposes to vasomotor syncope. A third type is anoxic and is seen in congenital heart disease with right to left shunt

Further details and treatment of the various forms of cardiac syncope are considered elsewhere

VASOMOTOR SYNCOPE

Under this heading may be grouped all varieties of syncope in which the cerebral blood flow fails as a result of a sudden fall in blood pressure due to collapse of the peripheral resistance. This includes the common faint

ETIOLOGY

Vasomotor syncope may be initiated by a critical fall in central venous pressure, by chemical agents that cause sudden profound vasodilatation or by stimulation of an assortment of receptors which excite a vaso-vagal reaction (Lewis 1932). Particular causes are listed below

Causing critical fall in central venous pressure

- 1 Haemorrhage
- 2 Loss of plasma into wounds burns crush injuries or gassed lungs
- 3 Loss of plasma into the skin or tissues as a result of allergy e.g. generalised urticaria and Quincke's oedema
- 4 Venous tourniquets on the thighs
- 5 Orthostatic hypotension
- 6 Other forms of postural hypotension

Chemical agents causing sudden profound vasodilatation

- 1 Acetylcholine and other cholinergic substances
- 2 Histamine
- 3 Tetraethylammonium salts
- 4 Nitrites

Stimulation of other receptors that excite a vaso-vagal reaction

- 1 Psychogenic disturbances
- 2 Carotid sinus compression
- 3 Extreme pain
- 4 Myocardial infarction
- 5 Pulmonary embolism
- 6 Meniere's syndrome

This list is by no means complete but it includes all the common causes of vasomotor syncope

MECHANISM

Syncope from haemorrhage has been thoroughly investigated in blood donors. As the blood volume diminishes the venous pressure falls and the cardiac output is reduced. Compensatory vasoconstriction may temporarily maintain the blood pressure. The faint which is associated with a sudden fall in blood pressure and pronounced bradycardia appears to be due to sudden vasodilatation in muscle (Barcroft *et al.* 1944). This vasodilatation is mediated by vasomotor nerves (Barcroft and Edholm 1944). Whether this reflex is excited by the fall in venous pressure or otherwise is unknown but it is clear that diminution in the blood volume is not directly responsible for the faint for the cardiac output may not alter at the critical moment—the peripheral resistance simply collapses.

This sequence of events has also been demonstrated when syncope results from the prolonged application of venous tourniquets to the thighs and probably occurs in all cases of syncope initiated by a critical fall in central venous pressure (Sharpey-Schafer 1944). Venous tourniquets on the thighs act as a bloodless venesection by trapping blood in the legs. Fainting in soldiers on parade who may have to stand at attention for long periods is believed to depend on similar factors. The fall in central venous pressure initiating orthostatic syncope following lumbo-dorsal sympathectomy is due

to abolition of veno motor tone in the lower half of the body. Veno motor paralysis may also be partly responsible for fainting following the injection of tetraethylammonium salts. Spontaneous, toxic, and convalescent orthostatic syncope may also be due to loss of veno motor tone.

Other forms of postural syncope include fainting in pregnant women when they lie on their backs too long and fainting in certain subjects on adopting the lordotic position. The fall in central venous pressure is then attributed to compression of the inferior vena cava by a pregnant uterus or by the liver which is forced against the spine (Bull 1948).

Syncope from chemical agents which cause sudden profound vasodilatation is directly due to collapse of the peripheral resistance. The blood pressure falls steeply but the cardiac output may be raised and there is usually tachycardia instead of bradycardia. Heat, gross aortic incompetence and other vasodilatation states predispose to syncope by lowering the peripheral resistance.

The simple psychogenic faint is initiated by emotional disturbance or by stimulation of the afferent component of a conditioned reflex; both result in a powerful autonomic discharge. The type of emotion usually responsible is a mixture of fear, amazement and curiosity, as may arise when a nurse sees a thoracic paracentesis for the first time or when a hypersensitive subject witnesses a street accident. The vasomotor centre appears to be suddenly depressed and there are associated cholinergic manifestations; the chief result is gross vasodilatation. This is certainly not in the skin which is pale and cold but may be in muscle or in the splanchnic bed. The peripheral resistance collapses and the blood pressure sinks rapidly. As the cerebral blood flow depends chiefly upon the blood pressure it becomes inadequate and consciousness is lost. Spontaneous recovery is inevitable for three reasons: first unconsciousness abolishes the trigger; secondly liberated acetylcholine upon which many of the features of the attack may depend is rapidly destroyed by choline esterase; thirdly the horizontal position naturally adopted by an unconscious subject increases the cardiac output and is favourable to the cerebral blood flow.

Carotid sinus syncope is said to be of four main types which may be reproduced by carotid sinus compression (Weiss and Baker 1933; Ferris, Capps and Weiss 1935). First syncope may be due to cardiac standstill (fig 601). Second loss of consciousness may be associated with a gross fall of blood

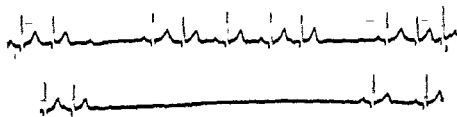


Fig 601—Carotid sinus pressure causing cardiac standstill

pressure and with marked slowing of the pulse rate. If the latter is restored to normal by atropine consciousness is not regained if the blood pressure is restored by any means consciousness returns even though the pulse remains slow. It is the low blood pressure and not the slow pulse rate which is responsible for the syncope. This type corresponds to vaso-vagal syncope. Third carotid sinus pressure may induce syncope associated with a profound fall in blood pressure without slowing of the pulse rate. It is doubtful if there is any fundamental difference between these two forms of attack for not infrequently the first type merges into the second indeed it has been suggested that initial slowing of the heart occurs in all cases but that subsequent quickening resulting reflexly from the low blood pressure may occur so rapidly as to mislead the observer.

Weiss and Baker describe a fourth type of syncope resulting from carotid sinus pressure in which the blood pressure and pulse rate are unchanged and refer to it as cerebral syncope. This appears to be allied to epilepsy for no reduction of cerebral blood flow can be demonstrated.

Spontaneous carotid sinus syncope may occur in rare instances. The organ is hypersensitive and may be excited by sudden pressure of the neck against a tight collar. The condition may be cured by carotid sinus denervation.

Reflex syncope from pain myocardial infarction pulmonary embolism etc. is similar in mechanism to the simple psychogenic faint.

CLINICAL FEATURES AND DIFFERENTIAL DIAGNOSIS

The chief clinical problem is the differentiation of vasomotor or vaso-vagal syncope from epilepsy. The difficulty lies in obtaining accurate data for the evidence in the first place is entirely historical physical examination is almost valueless as both are disorders of function not of structure. Further the statements of witnesses and not only lay witnesses are notoriously unreliable. In making a diagnosis reliance should be placed on information gained by thorough observation of a spontaneous or induced attack. Certain psychiatric or physical features may favour psychoneurosis on the one hand or epilepsy on the other but undue weight should not be attached to this.

The majority of epileptics have no warning whatsoever. If they have an aura it is odd and is not related to the autonomic nervous system. In sharp contrast vasomotor syncope is ushered in with numerous signs and symptoms of autonomic disturbance e.g. yawning pallor sweating coldness of the skin a sinking feeling in the pit of the stomach general muscular weakness subjective changes of temperature a feeling as if the blood was all rushing downwards epigastric discomfort and nausea desire to micturate or defæcate a feeling of light headedness or faintness and so forth.

The onset of epilepsy is neither quick nor sudden it is instantaneous at one moment the patient is in full possession of his senses a split second later he is unconscious. This means that he is unaware of the onset. He

may have a fit at night in bed and know nothing of it. Patients with vasomotor syncope on the other hand, feel themselves fainting. They lose consciousness gradually and although the onset may be described as quick or sudden, it is never abrupt.

The epileptic may have a fit at any time when he is walking, standing still, sitting lying or sleeping when he is in company or alone but rarely when his attention is concentrated. Psychoneurotics faint when standing up rarely when sitting and practically never when lying. They faint in company or when in reach of company rarely when alone. They are especially liable to attacks in closed spaces in church in the cinema and in circumstances that provoke emotional disturbance.

In epilepsy muscle tone is usually increased so that the patient falls rigid like a nine pin or if heightened tone is asymmetrical or local he may fall in a bent or twisted position. He is usually discovered lying prone so that he is apt to drown himself in shallow water or suffocate in his pillows. In contrast the muscles are flaccid in vasomotor syncope so that the patient collapses like a house of cards his final position being determined by gravity.

There are so many varieties of epilepsy that it is difficult to describe all the features which may occur during a fit but attention should be directed to increased muscle tone seen especially in the rigid phase of grand mal to clonic or regular jerking movements of hand trunk or limbs to conjugate deviation of the eyes to one side to Jacksonian localisation and to the epileptic march reflecting the anatomy of the motor cortex. Tonic contraction of the jaws and protrusion spasm of the tongue may result in the latter being bitten. Incontinence of urine may occur. If the central nervous system can be examined during an attack the eyes will be found open the pupils dilated and insensitive to light the corneal reflexes absent and the plantar response extensor.

During the rigid phase of grand mal breathing is impossible the subject becomes increasingly cyanosed the pulse rate and venous pressure rise and the heart pounds. During the physical effort associated with clonic convulsions the cardiovascular system behaves as it does with ordinary exertion.

In vasomotor syncope the patient lies flaccid and inert in a sprawled or crumpled position and may well be on his back. He is deathly white and often cold and clammy. The eyes may be open or closed the position of the upper lid being governed by gravity. The pupils are dilated and may be insensitive to light the reflexes and tendon jerks absent or depressed. The tongue is never bitten but urine may be voided. The essential feature is the low blood pressure which may be in the region of 50 or 60 mm Hg more often it cannot be determined. The pulse rate is slow normal or quick. In severe attacks slight twitching may be seen but is unimportant.

The epileptic attack is measured in seconds or minutes and lasts longer than three minutes. Vasomotor syncope is more variable and usually of short duration may last much longer even up to

Consciousness is regained as abruptly as it is lost in epilepsy it is regained gradually in vasomotor syncope

The epileptic may complain of headache and somnolence after an attack and of generalised muscle pains and aches after grand mal but if he does not pass off into a deep sleep and if he does not suffer from post epileptic automatism he recovers completely at once After vasomotor syncope the patient feels weak and ill he may complain of headache nausea or vomiting of a continued feeling of faintness or light headedness of trembling and shaking or of cold sweats He rarely recovers completely for half an hour or so and usually likes to lie down until he is better

Unfortunately both epilepsy and vasomotor syncope may be complicated by superimposed hysterical reactions which being more dramatic tend to impress both patient and witnesses to the exclusion of more vital phenomena This is one reason why historical diagnosis is often so difficult Pure hysterical fits have to be differentiated and this may not always be easy either by cross examination of the patient or of a witness Hysterical patients however never have an attack when alone never hurt themselves and may remember too much about it such as being aware of what is going on but being unable to move or lift a finger to help themselves Their movements during the period of alleged unconsciousness or trance like state do not fit in with the epileptic march the duration of the attack is usually longer and on recovery they do not complain of sore tongue sore muscles or of having urinated Despite all these points of difference if both patient and lay observer are bad witnesses the distinction between epilepsy hysteria and some other form of syncopal attack may be difficult In such cases electro encephalography may be helpful or the patient may be admitted to hospital for observation and if no spontaneous attacks occur one may be induced by the water pitressin test which depends upon the power of water retention to precipitate an attack of epilepsv

The patient is put on an ordinary diet and made to drink at least six pints (3.5 litres) of fluid daily When the body weight has increased by at least 3 lb (1.5 kg) which usually takes about 48 hours he is given 0.25 ml of pitressin intramuscularly and 300 ml of water by mouth thereafter he is given 0.5 ml of pitressin and 300 ml of water every two hours to a total of ten doses if necessary In epileptic subjects a fit is commonly produced after about the fifth or sixth injection and allows a correct diagnosis to be made in over 85 per cent of cases

Another simple diagnostic method is the hyperventilation test which may induce the patient to have one of his attacks

Vasomotor or vaso vagal syncope initiated by *haemorrhage* or by any other physical state which lowers the central venous pressure does not differ clinically from its psychogenic prototype Careful analysis of the circumstances under which the faint occurs may indicate the nature of the causal agent

Syncope may be produced by the *intravenous injection of acetylcholine*,

mecholin (acetyl beta methylcholine) and *doryl* (carbo amino acetylcholine) Loss of consciousness is preceded by flushing and a feeling of warmth due to vasodilatation and by sweating There is commonly abdominal colic nausea or vomiting and desire to micturate or defæcate The blood pressure is low but the pulse rate accelerates Patients may complain bitterly after regaining consciousness saying they feel dreadfully weak as if they had been ill for months Ordinary therapeutic doses of *mecholin* and *doryl* rarely cause syncope the dose must be large and given intravenously Symptoms are relieved at once by 1 to 2 mg. of atropine

The *intravenous injection of histamine* may also cause syncope due to gross general vasodilatation and collapse of the peripheral resistance Loss of consciousness is preceded by flushing and headache

An interesting and not uncommon form of syncope in elderly subjects may be closely associated with flushing It is encountered in menopausal women and occasionally in men at the climacteric Both flushes and fainting disappear following the administration of stilbæstrol 1 to 5 mg. daily

Meniere's syndrome or aural vertigo may occasion difficulty in diagnosis Vertigo is usually recognised by its spinning quality but occasionally there is no spinning but merely unsteadiness imbalance, or sudden attacks in which the subject is thrown violently forwards or backwards Consciousness is not lost however tinnitus is usually associated and deafness can nearly always be demonstrated

TREATMENT OF VASOMOTOR SYNCOPE

The causal agent should be identified and counteracted when possible Orthostatic hypotension may be improved by an abdominal binder or by bandaging the limbs Provocative postures should be avoided and patients should be instructed to stand up slowly Stilbæstrol may be tried in menopausal subjects

Psychogenic syncope calls for reassurance and psychotherapy for the fear of fainting encourages the faint Fortunately there is no danger attached to vasomotor syncope Patients usually have sufficient warning to ward off attacks by lying down or by sitting with the head between the knees Cool fresh air and bathing the face with cold water are helpful Brandy sal volatile and other stimulants may also be given or a glass of cold water may be preferred Half an hour on a couch in quiet comfortable and sympathetic surroundings consolidates recovery

CEREBRAL SYNCOPE

Cerebral syncope may result from cerebral vascular spasm or transient occlusion The fault is local

Hyperventilation syncope is the best example Forced breathing results in carbon dioxide washout with secondary tissue alkalosis Carbon dioxide ordinarily helps to maintain an adequate degree of cerebral vasodilatation

(Norcross 1938) its lack causes cerebral vasoconstriction. This induces dizziness within a minute in most normal individuals undergoing forced breathing. If hyperventilation is maintained long enough syncope may occur. Spontaneous attacks are seen in hysteria and sometimes in encephalitis lethargica. There is usually associated vasoconstriction in the extremities with pallor, cyanosis and tingling of the fingers and toes and there may be tetany. The blood pressure is maintained or raised owing to vasoconstriction, the latter tending to prevent reduction of cerebral blood flow.

Forced breathing may be used as a test in cases of syncope to discover whether an attack can be reproduced. It should be remembered, however, that epilepsy is sometimes excited by hyperventilation, so that the diagnosis depends upon the nature of the induced attack, not upon the simple fact that consciousness is lost. The effects of spontaneous hyperventilation may be quickly abolished by the inhalation of carbon dioxide. This may be accomplished by breathing in and out of a paper bag or long rubber tube.

Loss of consciousness due to hypertensive encephalopathy or to cerebral vascular lesions with or without associated spasm of cerebral vessels is usually called coma, unless convulsive epilepsy occurs. Embolism, however, especially when due to air or fat, may provoke an attack which fulfills the definition of syncope. The onset is abrupt and recovery may be remarkably quick and complete if the embolism moves on, or if spasm passes off suddenly.

Loss of consciousness occasionally occurs in Miniere's syndrome, but is then probably a vaso-vagal reaction.

Bilateral carotid compression, an old ju-jitsu trick, is a most effective way of inducing unconsciousness in an adversary.

ANOXIC SYNCOPE

Loss of consciousness resulting from most causes of anoxia is described as asphyxia or coma. Anoxic syncope, however, may occur in congenital heart disease with right to left shunt, when some factor suddenly reduces the amount of blood sent through the lungs; this factor may be something which increases the volume shunted, such as effort or screaming, or it may be something which reduces the venous return to the right auricle, such as an ill-advised venesection, or the sudden adoption of the upright posture by a bed-ridden patient.

Syncope in pilots used to be due to anoxia at high altitudes, but this has been overcome by the controlled use of oxygen. Nowadays it is chiefly associated with power dives and is governed by centrifugal forces. Syncope in anæmic subjects is usually vasomotor or vaso-vagal, anæmia merely acting as a predisposing factor.

REFERENCES

- Barcroft H and Edholm O G (1944) (Unpublished report to the Medical Research Council) ——— McMichael J and Sharpey Schafer E P (1944) Posthæmorrhagic fainting Study by cardiac output and forearm flow *Lancet* **i** 489
- Bull G M (1948) Personal communication
- Ferris E B Capps R B and Weiss S (1935) Carotid sinus syncope and its bearing on the mechanism of the unconscious state and convulsions *Medicine* **14** 377
- Lewis T (1932) Vaso vagal syncope and the carotid sinus mechanism *Brit med J* **i** 873
- Norcross N C (1938) Intra cerebral blood flow an experimental study *Arch Neurol and Psychiat* **40** 291
- Sharpey Schafer E P (1944) Circulatory dynamics of hæmorrhage *Brit med Bull* **2** 171
- Weiss S and Baker J P (1933) The carotid sinus reflex in health and disease Its role in the causation of fainting and convulsions *Medicine* **12** 297

CHAPTER VII

CONGENITAL HEART DISEASE

CONGENITAL anomalies account for 1 to 2 per cent of all cases of organic heart disease (Brown 1939) They are not hereditary and they are rarely familial The majority are due to defective development between the fifth and eighth week of foetal life some depend upon persistence of certain parts of the foetal circulation which should become obliterated at birth a few appear to be caused by infection *in utero* or are associated with German measles in the mother (Swan 1943) Other congenital abnormalities are found in at least 10 per cent especially arachnodactyly and mongolism Twins whether dizygotic or identical are rarely both affected

CLASSIFICATION

It has been customary to divide congenital heart disease into acyanotic and cyanotic forms and to subdivide the latter into types with permanent cyanosis (*morbis cœruleus* or blue babies) and types with late terminal or transient cyanosis (*cyanose tardive*) This has never proved entirely satisfactory and a new classification is therefore offered It is based on a series of 200 proved clinical cases (Wood 1950) and takes function into account

NO SHUNT		
GENERAL	LEFT SIDED	RIGHT SIDED
Dextrocardia Idiopathic hypertrophy Von Gierke's disease Heart Block Familial cardiomegaly	Coarctation of the aorta Right sided aortic arch Complete or incomplete aortic rings Bicuspid aortic valve (osup trunclar y cusp) Aortic sub stic tenion Left coronary artery arising from pulmonary artery	Idiopathic dilatation of the pulmonary artery Simple pulmonary stenosis Ebstein's disease

WITH SHUNT	
ACYANOTIC LEFT TO RIGHT SHUNT (pulmonary plethora)	CYANOTIC RIGHT TO LEFT SHUNT
<i>Left ventricular enlargement</i> Patent ductus Ventricular septal defect (with or without mild pulmonary stenosis) Perforated aortic sinus into P A or R V (or R A) Aorto pulmonary septal defect	DIMINISHED PULMONARY BLOOD FLOW LOW P A PRESSURE <ol style="list-style-type: none"> 1 <i>Left ventricular enlargement</i> Tricuspid atresia 2 <i>Right ventricular hypertrophy</i> Fallot's tetralogy Pulmonary atresia (Fallot type) Persistent truncus Pulmonary stenosis with reversed interatrial shunt
<i>Right ventricular enlargement</i> Atrial septal defect Anomalous pulmonary veins joining S V C or R A	HIGH P A PRESSURE Eisenmenger's complex Pulmonary hypertension with reversed aorto pulmonary, interventricular or interatrial shunt PULMONARY PLETHORA Transposition

DEXTROCARDIA

Mirror image dextrocardia is almost invariably associated with complete transposition of the viscera. The heart is functionally and structurally healthy. The electrocardiogram for obvious reasons shows reversal of all complexes in lead I with leads 2 and 3 interchanged (fig 7 01)

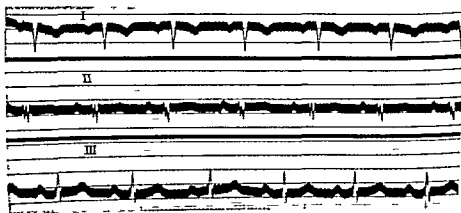


Fig 7 01—Electrocardiogram showing reversal of all complexes in lead I while leads 2 and 3 are interchanged

IDIOPATHIC HYPERTROPHY OF THE HEART

A rare condition not often compatible with more than a few years of life is that characterised by general enlargement of the heart progressing to early failure. Some of these cases have proved to be examples of Von Gierke's disease, the enlargement being due to glycogen retention in the muscle fibres of the heart as well as in the liver and other organs. There is acetonuria, a low fasting blood sugar, and a flat blood sugar curve following the injection of adrenaline, indicating failure of mobilisation of glycogen. Others are due to an anomalous origin of the left coronary artery from the pulmonary artery (Bland White and Garland 1933). Yet others prove to be cases of isolated myocarditis (Fiedler's carditis). There remains a group in which the etiology is obscure (Kugel 1939): gross thickening of the endocardium with embryonic myxomatous tissue richly supplied with elastic fibres (fibroelastosis) is usually found at necropsy, and may be the primary developmental defect (Glynn and Reinhold 1949).

The heart is grossly enlarged but remains normal in shape. There are no murmurs or other special features. The diagnosis depends on the finding of gross cardiac enlargement for no apparent reason. The prognosis is bad and there is no effective treatment.

FAMILIAL CARDIOMEGALY

From time to time cases of idiopathic cardiac enlargement are encountered in young subjects for which there is as yet no adequate explanation. X-rays show considerable cardiac enlargement, particularly of the left ventricle (fig. 702). Left bundle branch block is usually found. These patients are apt to die suddenly, presumably from ventricular fibrillation, or by degrees from congestive heart failure when still relatively young.

Some of these cases appear to have a familial basis (Addaru *et al.* 1946; Evans 1947). Necropsy reveals little but cardiac enlargement. Von Gierke's disease, isolated myo-



Fig. 702.—Unexplained cardiac enlargement in a relatively young man (there was also left bundle branch block).

carditis nutritional cardiopathies and abnormal coronary vessels must be excluded

COARCTATION OF THE AORTA

The word coarctation comes from the Latin *coarctatus* meaning pressed together tightened or contracted. As applied to the aorta it means a stricture of the arch usually just below the origin of the left subclavian artery.

Embryology. It will be recalled that there are two primitive aortas each having a ventral and a dorsal part joined by an arch. These three parts are called respectively the ventral aorta, the dorsal aorta and the first aortic arch. In front the two ventral aortas fuse to form a single tube from which develops the primitive heart the truncus arteriosus and the common ventral aorta. The two dorsal aortas also fuse between the fourth thoracic and fourth lumbar segments forming a single trunk the common dorsal or descending aorta. Caudal to the first pair of aortic arches spring five other pairs the six corresponding to the six branchial arches (fig 7 03a). In fishes these six vascular arches persist and supply the gills with blood for oxygenation.

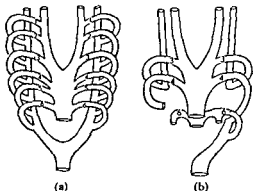


Fig 7 03 (a)—The six primitive aortic arches
(b) Subsequent arrangement of the six primitive aortic arches in man (see text)

In man and mammals subsequent development is illustrated in figure 7 03b. The first, second and fifth arches disappear. The third becomes the common carotid artery, the external carotid springing from it anteriorly, the internal linking up via the cranial portion of the dorsal aorta. The fourth arch

becomes the proximal part of the subclavian artery on the right side and the final aortic arch proximal to the junction of the ductus arteriosus on the left. The sixth arches are separated from the aortic system by the aorto pulmonary septum which divides the truncus into anterior and posterior halves; the anterior half becomes the ascending aorta, the posterior the pulmonary artery. The division of the truncus extends cranially to a point just beyond the anterior ends of the sixth arches, the mouths of which are included in the posterior section and therefore in the pulmonary system. On the right side the sixth arch becomes the right pulmonary artery and loses its connexion with the right dorsal aorta; on the left it becomes the left pulmonary artery and preserves its connexion with the left dorsal aorta in the form of the ductus arteriosus. While these changes are going on harmonious alterations take place in the ventral and dorsal aortas. In front

the two ventral aortas fuse into a single ascending aorta as already indicated. Behind the dorsal aortas undergo considerable modification the upper part forms a portion of the internal carotid artery as previously described the segment between the third and fourth arches disappears caudal to the fourth arch the dorsal aorta disappears on the right side except for that part of it which is incorporated in the right subclavian artery and forms the posterior part of the aortic arch on the left side. The left subclavian artery links up with the left dorsal aorta just below the junction of the sixth arch i.e. just below the ductus.

Many anomalies may result from faulty development of this aortic system. Thus the caudal part of the right dorsal aorta may persist so that there are two aortic arches or the caudal part of the left dorsal aorta may disappear in favour of the right so that the final aortic arch is right sided. The most important however is partial obliteration of that part of the left dorsal aorta which lies between the fourth and sixth arches i.e. just above the ductus or between the sixth arch and the point of fusion of the two dorsal aortas i.e. just below the ductus. This short segment of the aorta is

often called the isthmus on account of the frequency with which it is narrowed but in coarctation or isthmus stenosis narrowing is extreme and often remarkably abrupt. There are two main types infantile and adult (Bonnet 1903). In the former (fig 7.04a) the constriction is above the ductus which remains patent and carries venous blood to the descending aorta being incompatible with more than a few years of life it will not be further con-

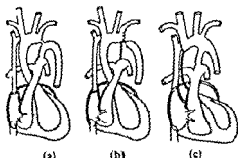


Fig 7.04—Diagram illustrating the three main types of coarctation of the aorta

(a) Infantile type with patent ductus feeding the descending aorta

(b) Common adult type

(c) Aortic atresia with patent ductus feeding the whole systemic circulation

sidered. In the latter (fig 7.04b) the ductus is closed or if patent the constriction is below it so that it plays no part in compensating for the defect. Aortic atresia with a patent ductus feeding the whole systemic circulation (fig 7.04c) constitutes a third type (Bramwell 1947) but such cases all die in infancy. Other variants of these three main types have been described by Evans (1933).

Hemodynamics. The clinical features of the adult form of coarctation depend upon the mechanical effect of the constriction and upon the development of an extensive collateral circulation. Much use is made of the branches of the subclavian artery e.g. the superior intercostal and the internal mammary with its intercostal superior epigastric and musculo-phrenic rami also of the thoracic and subscapular branches of the

axillary artery. These vessels link up with the intercostal branches of the descending aorta and with the inferior epigastric branches of the femoral arteries and so by-pass the constriction. The blood pressure is elevated in vessels arising from the aorta above the isthmus below it the systolic pressure is reduced and the diastolic raised the mean pressure often being still above normal. The cause of the hypertension is uncertain. The raised mean pressure in the legs does not support the mechanical hypothesis. Renal ischaemia was blamed by Ryland (1938) on the grounds that hypertension was only produced experimentally when the aorta was constricted above the origin of the renal arteries but the renal blood flow in clinical cases of coarctation appears to be normal. The truth may not be discovered until the cause of essential hypertension is known.

CLINICAL FEATURES

Coarctation occurred in 8 per cent of the author's series of 200 cases of congenital heart disease. It is said to be 4 to 5 times more common in men than in women but in the author's series the sex incidence was equal. Hypertension is usually considerable in degree and may be discovered in childhood. Patients may complain of nose bleeding or of undue throbbing. All the usual features of hypertensive heart disease are present and this is responsible for death from congestive failure in about 40 per cent of cases (Abbott 1928). Renal failure does not occur because the kidneys are protected by the lesion but cerebral accidents are not uncommon. Other symptoms include gnawing pains in the shoulder girdle, chest or back, cold feet and occasionally intermittent claudication.

The major peripheral arteries should be palpated as a routine in all cases of hypertension. In coarctation femoral pulsation is not only feeble but delayed and pulsation in the posterior tibial and dorsal arteries of the feet may be absent. If the blood pressure in the legs can be measured at all by the cuff method it will be found to be lower than in the arms but if measured directly by means of a needle in the femoral artery and an electrical manometer the diastolic pressure is found to be raised though the systolic is reduced in other words the blood pressure in the legs approaches the mean pressure the amplitude of pulsation being diminished (Brown *et al* 1948).

A mitral diastolic murmur is by no means uncommon even in early childhood and is usually attributed to coincident rheumatic endocarditis or to established mitral stenosis. There is no evidence as yet that such a murmur is ever functional in coarctation.

The collateral circulation provides the majority of the physical signs enlarged tortuous vessels giving rise to unusual pulsations and murmurs. Pulsation of some of the intercostal arteries can usually be seen and felt posteriorly especially if the subject bends forward and may be associated with similar murmurs. Radiography may reveal notching of the inferior borders of the ribs (fig 7 o₃) due to pressure erosion (Railsbach and Dock

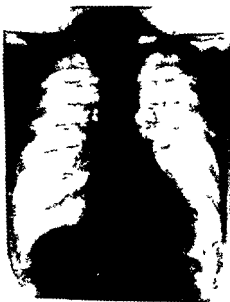


Fig 7.05—Notching of the inferior border of the ribs due to pressure erosion from enlarged intercostal arteries in coarctation of the aorta. The aortic knuckle is elongated and is formed by a grossly enlarged left subclavian artery.



Fig 7.06—Visualisation of coarctation of the aorta by means of angiocardiology (Courtesy of Dr Wallace B. Eden).

19.9) this may be seen in children as young as six but is more obvious in adults.

The constriction itself is difficult to see fluoroscopically but may be demonstrated clearly by means of angiocardiology (fig 7.06) (Grishman, Steinberg and Sussman 1941) or retrograde aortography (Broden, Hanson and Karnell 1948). Hypoplasia of the aortic knuckle as seen in the ordinary antero-posterior skiagram is suggestive especially when associated with left ventricular enlargement. A double aortic knuckle representing the blind ends of the aorta above and below the constriction is diagnostic (Bramwell 1947); it is usual for the upper knuckle to be elongated being formed by a grossly enlarged left subclavian artery (fig 7.05).

ASSOCIATED PHENOMENA

Bicuspid aortic valves may be associated (11 per cent in Abbott's series (1928), 42.3 per cent in that of Reifenshtein *et al.* (1947)) and may lead to *aortic incompetence* under the stress of the hypertension. *Patent ductus arteriosus* with the usual clinical features occurs in 10 per cent of adult coarctations (Bramwell 1947). Owing to the high pressure gradient between the arch of the aorta and the pulmonary artery the blood flow through the

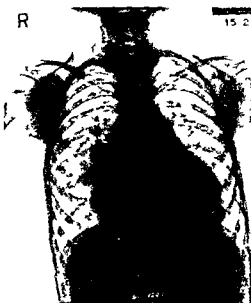


Fig 7 07 (a)—Coarctation of the aorta associated with patent interventricular septum proved at necropsy

Fig 7 07 (b)—Similar case but without necropsy proof

ductus may be enormous the pulmonary vessels are thus unusually dense and dilated the left auricle may resemble that in mitral stenosis and there is often a mitral diastolic murmur At the same time the systemic collateral circulation may be less in evidence owing to the ready escape of aortic blood into the pulmonary circulation

A very similar picture may be seen when coarctation is associated with patent interventricular septum the high pressure in the left ventricle causing an unusually large left to right shunt In a case investigated by the author and subsequently proved at necropsy the mean pulmonary arterial pressure was 130 cm of saline whilst that in the right ventricle was 65 to 82 Samples from the pulmonary artery were 86 per cent saturated with oxygen from the middle of the right ventricle 70 per cent and from the right auricle and superior vena cava 60 per cent Clinically coarctation of the aorta was recognised by the presence of high blood pressure in the carotid and subclavian arteries (160/100 mm Hg in a boy of six) with an immeasurable pressure in the legs but there was little evidence of a collateral circulation The pulmonary arteries were grossly engorged radiologically (fig 7 07a) there was a pulmonary diastolic murmur at the base and a mitral diastolic murmur with triple rhythm at the apex Despite the absence of a machinery murmur patent ductus arteriosus was believed to be responsible for the shunt and seemed to be confirmed by the catheter

findings the raised oxygen content of the right ventricular sample being attributed to pulmonary incompetence. At necropsy coarctation of the aorta of the adult type was associated with a large defect of the membranous interventricular septum. The aortic cusps were normal the aortic ring admitted only the little finger and the ascending aorta was small. The defect in the septum admitted the middle finger whilst the pulmonary ring admitted both middle and fore fingers. A very small patent ductus joined the aorta below the isthmus. Although the huge pulmonary artery did not sit astride the septal defect there could be no doubt that the major portion of the left ventricular contents was expelled into that vessel. The mitral diastolic murmur was clearly functional for there was no sign of mitral stenosis. Both ventricles were greatly enlarged the left retaining its natural dominance. A second example in a man with precisely the same clinical features is illustrated in fig 7 07b. Cardiac catheterisation excluded atrial septal defect but did not distinguish patent ductus with pulmonary incompetence from interventricular septal defect. The patient died later from congestive heart failure but there was no necropsy.

Bacterial endarteritis may affect the isthmus or an associated bicuspid aortic valve. It develops in about 20 per cent of cases. *Berry aneurysm* may occur in the region of the circle of Willis and may result in *subarachnoid hæmorrhage* especially in those with higher blood pressures. Death from subarachnoid hæmorrhage occurred in 9 per cent of 200 cases reported by Maud Abbott (1928). Sometimes there is no aneurysm but the media or elastic tissue is defective (Glynn 1940). Similar defects may be found in the aorta and may be responsible for the frequency with which it ruptures (19 per cent of Abbott's series). *Variations in one or other subclavian artery* occur in about 5 per cent of cases (King 1937) and may be due to its anomalous (East 1932) or stenotic (I owe and Holms 1939) origin.

PROGNOSIS AND TREATMENT

Although many patients live well into middle life without serious handicap some even to the eighth decade the majority succumb between the ages of 20 and 40 to one of the complications mentioned above (Abbott 1928). The average age of death is 35 (Reifenstein Levine and Gross 1947). Surgical repair (Crafoord and Nylén 1945) should therefore be offered. The physiological results of such an operation are usually good the blood pressure falls symptoms disappear the heart becomes smaller and it may be assumed that the risks of intracranial hæmorrhage aortic rupture and bacterial endocarditis are diminished. Crafoord (1948) had successfully repaired 22 cases with only two deaths up to July 1947. The ages of his patients ranged from 11 to 27 but late childhood (age 8 to 12) may be the best time for the operation. Of 52 cases operated on by Gross (1949) there were 7 deaths 4 failures and 41 cures in this series there were eight additional cases that were considered inoperable. A mortality rate of 16 per cent amongst 128 surgically treated cases in different centres

was reported by Shapiro (1949) The chief dangers appear to be sudden cardiac asystole when the aorta is clamped especially if a large left subclavian artery has to be included in the clamp and tearing out of sutures from a hypoplastic aorta distal to the constriction Pre operative preparation and choice of anæsthetic designed to prevent an undue rise of blood pressure when the aorta is clamped may help to reduce the first risk

BICUSPID AORTIC VALVE

Bicuspid aortic valve cannot be diagnosed clinically without aortic incompetence or superimposed bacterial infection The former is unusual unless there is hypertension or infection for a healthy bicuspid valve may be competent against a normal blood pressure Bacterial endocarditis attacks about one-quarter of all cases

AORTIC STENOSIS

Aortic valvular stenosis is rare It differs little from acquired aortic stenosis but may be accompanied by stunted growth and weak physical development The majority die young

Subaortic stenosis is perhaps less rare (2 per cent) The lesion which affects the outflow tract of the left ventricle is due to defective absorption of the primitive bulbus cordis A perforated membrane lies proximal to the valve The clinical features differ from those of valvular stenosis in that the aortic second sound is clear the peripheral pulse usually normal the left ventricle but little enlarged and the prognosis apart from the risk of bacterial endocarditis excellent The diagnosis is not excluded by the deposition of calcium Some of these cases are mistaken for the *Maladie de Roger*

Both forms are more common in males than in females

IDIOPATHIC DILATATION OF THE PULMONARY ARTERY

Cases are seen occasionally with considerable dilatation of the pulmonary artery without obvious cause (Laubry Routier and de Balsac 1941) If there is functional pulmonary incompetence the right ventricle enlarges and partial or complete right bundle branch block may follow as in atrial septal defect The pressures in the right side of the heart are normal and the pulmonary blood flow is not increased The X ray appearances differ from those of ASD in showing normal pulmonary vessels beyond the two main branches The prognosis is relatively good

ATRIAL SEPTAL DEFECT

Embryology Atrial septal defect refers to a relatively large non valvular opening in the atrial septum through which blood may flow either way. Embryologically the atrial septum is formed in the first place by the sickle shaped septum primum which grows forwards from the dorsal wall of the common auricle dividing it into two. For a time communication exists between the two auricles in front of the crescentic edge of the growing septum. If development is arrested at this stage a septal defect results and is situated in the lower anterior part of the septum just below and usually including part of the fossa ovalis. When growth proceeds normally this hole is obliterated and a new one the foramen ovale appears in the upper and dorsal part of the septum primum. Arrest at this stage results in a defect just above the site of the fossa ovalis. With further normal development the foramen ovale comes to lie more anteriorly and is turned into a valve by the growth of the septum secundum on the right side of the septum primum and covering it at all points except over the area known as the fossa ovalis. When the septum secundum develops fully and the septum primum degenerates completely the defect occurs at the site of the fossa ovalis.

In *patent foramen ovale* the septa are fully developed but imperfectly fused. When pressure is applied to the right side of the fossa ovalis the septa are parted, blood penetrates between them and escapes into the left auricle through the patency in the upper part of the septum primum known as the foramen ovale proper. In foetal life the relatively high pressure in the right auricle keeps the valve open and causes blood to be shunted from right to left in order to avoid the pulmonary circulation. At birth the pressure rises in the left auricle and forces the septum primum against the septum secundum thereby closing the valve. In 80 per cent of all individuals fusion then takes place between the two septa and the foramen ovale is permanently closed. In the remaining 20 per cent fusion fails and valvular patency continues. Occasionally it causes terminal cyanosis in conditions such as pulmonary heart disease in which the right auricular pressure may come to exceed the left.

A cardiac catheter may slip through a patent foramen ovale into the left auricle without difficulty and may enter the left ventricle (fig 7 o8a) or any of the pulmonary veins (fig 7 o8b). The pressures and electrical potentials in these chambers may thus be obtained in favourable cases including otherwise normal hearts. The mean left auricular pressure appears to be 3 to 5 cm. of saline above the sternal angle and is distinctly higher than the right. Pulmonary venous samples have always been about 95 per cent saturated with oxygen whatever the patient's condition (this includes Fallot's tetralogy and other cases of cyanotic congenital heart disease but not anoxic pulmonary heart disease). Uncomplicated patent foramen ovale is easily distinguished from atrial septal defect because of the absence of any appreciable inter auricular shunt as judged by samples from both auricles and their respective venous systems.



Fig 7 08 (a)—Catheter in the left ventricle via a patent foramen ovale



Fig 7 08 (b)—Patent foramen ovale proved by cardiac catheterization the catheter has entered a pulmonary vein and has been pulled taut. The sharp angle towards the end of the catheter represents the foramen ovale



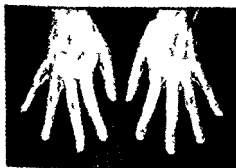
Fig 7 09—Catheter in left atrial appendage



(a) Facies (note bilateral arideotomy)



(b) Showing high arched palate and deformed teeth



(c) Spider fingers

Fig 7 to—A case of arachnodactyly

This patient was 6 ft high and also showed hypotonia scoliosis and flat feet

Another interesting anatomical study connected with this work has been the location of the left atrial appendix which has always been on the left border of the heart between the pulmonary arc and left ventricle (fig 7 09)

Hæmodynamics The defect in the septum is usually 1 to 3 cm in diameter and carries a considerable shunt from left to right auricle the pressure being higher on the left Oxygenated blood is thus added to the normal intake of the right ventricle the stroke output of which is correspondingly increased The situation is met by right ventricular enlargement and dilatation of the pulmonary artery Left auricular leakage deprives the left ventricle of its full intake the left ventricular stroke output is diminished the left ventricle and aorta hypoplastic and the pulse small Progressive right ventricular enlargement eventually leads to failure the pressure in the right auricle then rises and if it exceeds that in the left the shunt is reversed and cyanosis develops Acquired pulmonary hypertension may have a similar effect

Incidence A S D accounted for 17 per cent of the author's unselected series of 200 proved cases of congenital heart disease It shows a strong preference for females the sex ratio being 4 : 1 in their favour Associated mitral stenosis (Lutembacher's syndrome) is remarkably common especially in females being present in about 33 per cent of all cases Other valves may also be affected and adhesive pericarditis is not infrequent It may be significant that cases of arachnodactyly which appears to bear some relationship to atrial septal defect are also prone to develop rheumatic manifestations

Arachnodactyly (fig 7 10) is an hereditary and familial disorder of mesoblastic growth and is characterised by elongation of the fingers and toes thin facies tall lean build hypotonia (and its consequences) dislocation of the lenses high arched palate and pigeon chest Cardiac abnormalities especially atrial septal defect are associated in 40 to 45 per cent of cases

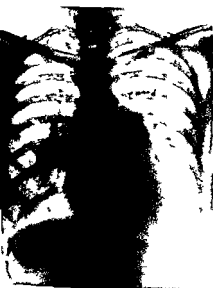
CLINICAL FEATURES

There may be no symptoms until the third or fourth decade effort tolerance being good and cyanosis absent hæmoptysis did not occur in 6 cases analysed recently at the Institute of Cardiology Between the ages of 25 to 45 however congestive failure usually develops and may be associated with the sudden appearance of central cyanosis

Physical signs include a small or normal peripheral pulse a normal jugular venous pressure without a conspicuous a wave visible or palpable pulsation of the pulmonary artery and outflow tract of the right ventricle a pulmonary systolic murmur with or without thrill and a widely split second sound at the base without accentuation of the second or pulmonary element not infrequently there is also a basal diastolic murmur due to functional pulmonary incompetence (Graham Steell murmur) At the apex beat only the tap of mitral valve closure may be discerned but



Fig. 7 11—Skadiogram of a case of atrial septal defect showing gross dilatation of the pulmonary artery and its branches, enlargement of the right auricle and hypoplasia of the aorta



(a) Anteroposterior view

(b) First oblique position showing dilatation of the right auricle

Fig 7 12—I sternbacher syndrome

occasionally the cardiac impulse is tumultuous being formed by the distended right ventricle



Fig 7 13—Atrial septal defect in a child aged 10

Fluoroscopy in well developed cases (fig 7 11) reveals gross dilatation and conspicuous pulsation (hilar dance) of the pulmonary artery and its branches considerable enlargement of the right auricle and ventricle hypoplasia of the aorta and left ventricle and a flat left auricle In Lutembacher's syndrome (fig 7 12) the left auricle is also enlarged and the right more so In less advanced cases however and especially in children the changes described may be much less noticeable (fig 7 13)

I lectrocardiograms show partial or complete right bundle branch block in 95 per cent of cases (fig 7 14) Auricular fibrillation occurs in 10 to 20

per cent and is usually due to associated mitral stenosis The extraordinary frequency of right bundle branch block is interesting in view of the relatively low pressure in the right ventricle, and suggests that dilatation of that chamber is responsible The wide splitting of the second heart sound may well be due at least in part to the conduction defect

The diagnosis may be proved by obtaining samples of relatively oxygenated blood from the right auricle right ventricle and pulmonary artery by means of cardiac catheterisation when samples from the *venæ cavae* show ordinary venous blood (Howarth McMichael and Sharpey Schafer, 1947) In twenty five cases investigated by the author (fig 7 15) samples obtained from the right auricle right ventricle and pulmonary artery differed little and ranged between 81 and 90 per cent saturated with oxygen caval samples being normal (62 to 73 per cent saturated) Samples from the pulmonary veins left auricle left ventricle and femoral artery were always between 91 and 96 per cent saturated Mean pressures in the pulmonary artery were lower than expected all but one ranging between 6 and 18 mm Hg above the sternal angle The mean left auricular pressure was distinctly raised and was 10 cm of saline in one case presumably because of the increased quantity of blood in the lungs This suggests that an atrial septal defect initiates a vicious circle for the shunt apparently increases the pressure gradient between the two auricles—as long as the

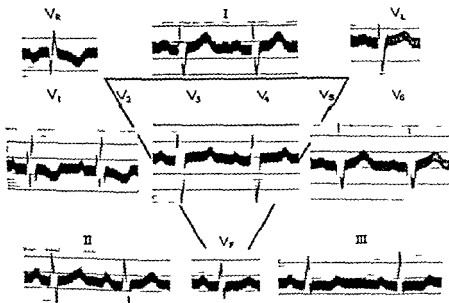


Fig 7:4 —Electrocardiogram in a case of atrial septal defect showing right bundle branch block

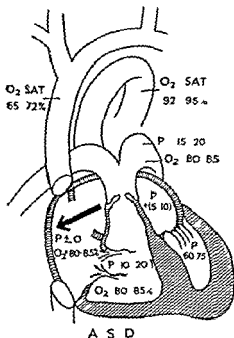


Fig 7:5—Functional studies in atrial septal defect (P=mean pressure in cm of saline above the sternum angle)

right ventricle functions efficiently. Simple calculations reveal that the pulmonary blood flow is usually two to three times the systemic flow and is of the order of 10 to 15 litres per minute.

Angiograms are less helpful but an artificial shunt from right to left auricle may be demonstrated sometimes owing to the sudden rise of right auricular pressure resulting from the large bulk of fluid injected so quickly and forcefully.

PROGNOSIS AND TREATMENT

Up to the age of 25 or so there is little disability but the situation may change radically during the fourth decade when heart failure (and perhaps permanent cyanosis) may develop. Bacterial endocarditis is extremely rare. The average age of death is 35 to 36 with or without mitral stenosis (McGinn and White 1933; Roesler 1934) but several cases over 70 have been recorded.

Restriction of effort is advisable even when there are no symptoms for the right ventricle must be spared unnecessary work. Should permanent central cyanosis develop digitalis and mercurial diuretics should be given whether or not there is auricular fibrillation and whether or not signs of systemic congestion are apparent for heart failure must be inferred (unless there is tricuspid incompetence).

Surgical repair is in the experimental stage (Murray 1948).

VENTRICULAR SEPTAL DEFECT

Ventricular septal defect commonly refers to an isolated defect of the membranous part of the interventricular septum due to failure of the aortic septum to fuse with the ventricular septum. Diagnosed clinically for the first time by Roger (1879) the lesion has been said to account for 35 to 37 per cent of all cases of congenital heart disease recognised at school age (Perry 1931; Muir and Brown 1934). Such high figures probably include instances of subaortic stenosis, simple pulmonary stenosis, mitral incompetence and innocent parasternal murmur.

In the author's series of 200 proved clinical cases of congenital heart disease isolated VSD occurred in 12 per cent and VSD with simple pulmonary stenosis in an additional 2 per cent.

HÆMODYNAMICS

Functional studies by the author in 16 unselected cases of VSD have shown mean pulmonary artery pressures of 7 to 95 mm of Hg the pressure being proportional to the shunt and to the peripheral pulmonary resistance. The latter was increased in three and reduced in four. True pulmonary hypertension from an increased resistance was not due to the size or duration of the shunt but to some inherent or predetermined factor. Samples from the right auricle have been normal (58 to 76 per cent

saturated with oxygen) At a low level in the right ventricle just proximal to the pulmonary valve and in the main trunk of the pulmonary artery saturation figures were similar and were in the region of 81 to 86 per cent saturated

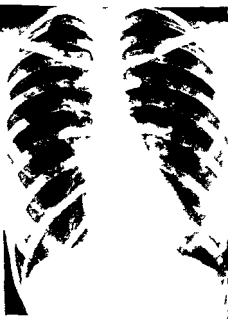


Fig 7 16—Maladie de Roger—X ray appearances



Fig 7 17—Skogram of a case of ventricular septal defect with considerable increase of pulmonary blood flow

The degree of shunt from left to right ventricle may be calculated by finding the difference between the blood flow through the pulmonary artery and that through the tricuspid valve

$$\text{Shunt flow} = \text{pulmonary flow} - \text{tricuspid (systemic) flow}$$

$$= \frac{O}{a-p} - \frac{O}{a-v}$$

where O is the oxygen consumption in ml per minute

a is the oxygen content of arterial blood

p is the oxygen content of pulmonary artery blood

v is the oxygen content of mixed venous blood in the right auricle

In the case mentioned above the pulmonary blood flow ranged between 1½ and 4 times the systemic flow and was usually between 11 and 16 L/min

CLINICAL FEATURES

The majority have no symptoms but severe cases may develop congestive failure in childhood or adolescence. There is no cyanosis.

The pulse is small or normal and the jugular venous pressure normal or slightly raised. The cardiac impulse is left ventricular in type and usually hyperdynamic. The right ventricular outflow tract is inclined to be lifting and pulmonary artery pulsation may be visible or palpable. A systolic thrill and murmur are nearly always present and are commonly maximal in the third and fourth intercostal spaces at the left sternal border but may be higher. A functional mitral diastolic murmur due to a torrential mitral blood flow is heard in at least half the cases and in nearly all severe cases. The second heart sound at the pulmonary area is normally split, the second or pulmonary element being commonly accentuated. A functional pulmonary diastolic murmur occurs in about a quarter of the more severe cases.

The electrocardiogram is normal in mild cases but shows prominent Q waves and tall R waves (with or without T or U wave changes) in left ventricular surface leads and conspicuous secondary R waves without much widening of QRS in leads V₁ or V₂ in the majority appearances suggesting both left and right ventricular enlargement.

X rays reveal pulmonary plethora, a varying degree of dilatation of the pulmonary artery and left ventricular enlargement in all but the mildest cases (figs 7.16 and 7.17) appearances that are indistinguishable from those of patent ductus.

It will be appreciated that these clinical features are not those described by Roger. It is suggested therefore that the term *Maladie de Roger* should apply only to mild cases which show nothing abnormal apart from the Roger thrill and murmur, i.e. to about one quarter or at most one third of all cases of VSD.

In differential diagnosis the *maladie de Roger* must be distinguished from innocent parasternal murmur, subaortic stenosis, mild pulmonary stenosis and organic mitral incompetence. VSD in its wider sense is more likely to be confused with atrial septal defect or patent ductus arteriosus. If due regard is paid to the quality of the cardiac impulse, to the nature of the pulse and to the electrocardiogram in addition to the site and character of the murmur and to the X ray appearances, a correct clinical diagnosis is usually possible.

Cases that develop true pulmonary hypertension are in danger of reversing the shunt; should this occur the situation would be clinically indistinguishable from the Eisenmenger complex (page 244).

There have been some interesting cases in which characteristic signs of a ventricular septal defect discovered in childhood have disappeared with advancing years. Whilst it is difficult to prove that these were not examples of innocent left parasternal murmur, it has been suggested that spontaneous

obliteration of small defects may sometimes occur (Parkes Weber 1918)

The prognosis would be good in mild cases were it not for the 25 per cent risk of bacterial endocarditis. When this occurs emboli are confined to the pulmonary circulation (unless infection spreads to the aortic valve) vegetations occurring round the defect on the right ventricular side and on the opposite wall of the right ventricle where the shunted blood stream impinges. Patients with pulmonary plethora are in a different category and may die from congestive heart failure in adolescence (Baldwin Moore and Noble 1946)

No reparative treatment is yet generally available but Murray (1948) has made the attempt. Mild cases should be encouraged to lead normal unrestricted lives but severe cases need care and should limit their physical activities. Dental treatment, sore throat and other pyogenic infections should be covered by a short course of penicillin to prevent endocarditis.

PATENT DUCTUS ARTERIOSUS

Incidence. Whilst failure of the ductus arteriosus to close within a few months of birth is often associated with other congenital anomalies it is relatively common alone especially in females the sex incidence being 2 : 1 in their favour (Benn 1947). It was the main or sole lesion in 9.2 per cent of 1,000 cases of congenital heart disease collected by Abbott and accounted for 14.5 per cent of the author's series.

Hæmodynamics. The ductus connects the left branch of the pulmonary artery with the arch of the aorta just opposite the origin of the left subclavian artery. As the aortic pressure is higher the blood flow is from aorta to pulmonary artery and its degree depends not only upon the size of the ductus but also upon the pressure gradient between these two vessels. There is thus an aortic leak the equivalent of an arteriovenous shunt and a raised pressure and blood flow in the pulmonary artery. The volume of blood entering the left auricle is greater than normal and the left ventricular stroke and minute output are increased. The total blood volume is also increased (Cassels and Morse 1947).

CLINICAL FEATURES

There are usually no symptoms when the lesion is first discovered.

Owing to the large stroke volume and aortic leak the pulse is abrupt and collapsing, the pulse pressure raised and the diastolic blood pressure rather low especially after exertion when it may drop grossly (Bohn 1938). The cardiac impulse is forceful, displaced to the left and suggests left ventricular enlargement. The classical machinery murmur (Gibson 1900) usually accompanied by a thrill begins just after the first sound, waxes towards the end of systole and wanes in late diastole. It is more or less localised to the second left intercostal space. Occasionally especially in infants and young children the murmur is confined to systole, very rarely it is entirely diastolic possibly due to pulmonary incompetence. If not

heard at rest it may be encouraged by pressor agents exercise amyl nitrite or Muller's experiment It may be reduced or abolished by raising the pulmonary arterial pressure or lowering the aortic e.g. by Valsalva's experiment breath holding making the patient breathe 10 per cent oxygen or by hypotensive agents The increased pressure within the pulmonary artery causes accentuation of the pulmonary element of the second sound In severe cases a functional mitral diastolic murmur is usually heard due to dilatation and rapid filling of the left ventricle—hence the not uncommon mistake of confusing patent ductus with aortic incompetence and mitral stenosis

Fluoroscopy (figs 7 18 and 7 19) reveals abrupt dilatation and conspicuous pulsation of the pulmonary artery pulmonary plethora enlargement of the left ventricle and fullness of the left auricle (Donovan Neuhauser and Sosman 1943) In contrast to atrial septal defect and mitral stenosis but like hyperkinetic pulmonary heart disease the aortic knuckle is normal or prominent A local bulge in the region of the aortic isthmus can be demonstrated by means of angiocardiology in a

limited number of cases and is thought to represent the widened mouth of the ductus or possibly a traction aneurysm of the aorta (Steinberg Grishman and Sussman 1943) With suitable technique angiocardiology may also show the pulmonary artery filling twice first from the right ventricle then from the aorta Retrograde aortography offers an alternative means of obtaining good angiograms

The electrocardiogram is usually normal but when the ductus is large there may be evidence of left ventricular hypertrophy (fig 7 20) Extreme cases may show the classical electrocardiographic changes of gross left ventricular enlargement (fig 7 21)

Fig 7 18—Diagram of a case of patent ductus showing enlargement of the left ventricle but little dilatation of the pulmonary artery



The diagnosis may be proved in doubtful cases by means of cardiac catheterisation (fig 7 22) Samples of blood from the superior vena cava right auricle and right ventricle are normal (about 70 per cent saturated) whereas samples from the pulmonary artery are usually 80 to 85 per cent saturated Slight admixture of arterial blood in right ventricular samples



Fig 7 19--Skiagram of a more advanced case of patent ductus showing considerable left ventricular enlargement and enlargement of the pulmonary vessels in addition to dilatation of the pulmonary artery

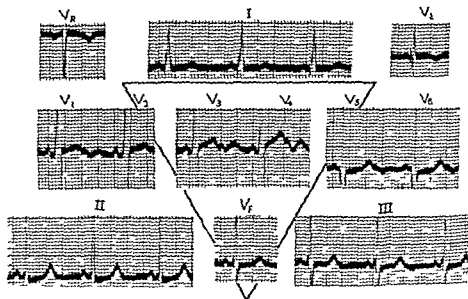


Fig 7 20—Electrocardiogram in a case of patent ductus showing left ventricular enlargement. There is a strong QR pattern with inverted U waves in lead V₅.

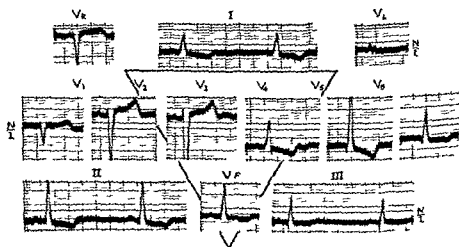


Fig 7 21—Cross left ventricular preponderance in a case of patent ductus.

should be left alone. Bacterial endocarditis should be cured by means of penicillin if possible before submitting the patient to operation.

The chief diagnostic errors that have resulted in fruitless thoracotomy have been due to mistaking some other source of a continuous murmur for a ductus, a normal venous hum at the base of the neck in a child, arterio-venous aneurysm of the lung, and perforation of the sinus of Valsalva into the pulmonary artery or right ventricle may be responsible for such a murmur. In cyanosed cases a machinery murmur may be due to direct communications between bronchial and pulmonary arteries.

Although pulmonary engorgement diminishes and the heart decreases in size after successful ligation in most instances (fig. 7.23) irreversible changes are encountered occasionally and for this reason severe cases should not be left too long.

PULMONARY STENOSIS

Pathogenesis. Subvalvular stenosis is thought to be due to arrested evolution of the bulbus cordis, most of which should become incorporated in the right ventricle (Keith, 1909). The obstruction is in the outflow tract or conus of the right ventricle. There are two kinds: in one the primitive bulbus forms a separate chamber between the pulmonary artery and right ventricle and communicates with the latter by a small lower bulbar orifice which is the cause of the stenosis; in the other the conus is diffusely constricted, leaving a narrow passage between the right ventricle and pulmonary valve. Valvular stenosis is proving more common than previously thought; the valve is then represented by a conical membrane with a small circular hole in the centre, or the cusps may be fused in a manner resembling inflammatory stenosis.

CLINICAL FEATURES

Simple acyanotic pulmonary stenosis with closed septa accounted for 10 per cent of the author's series of 200 cases of congenital heart disease. The stenosis is usually valvular, and cases are acyanotic until reduction of cardiac output and compensatory peripheral vasoconstriction lead to a sluggish surface blood flow. In other words, cyanosis, when it occurs, is peripheral, not central, the arterial oxygen saturation being normal. The chief symptoms are progressive breathlessness on exertion and ultimately those of congestive failure. In severe cases angina pectoris and frightening syncopal attacks may occur.

In relatively severe cases the physical signs include a conspicuous *a* wave in the jugular pulse (which may be transmitted to the liver), a high pulmonary systolic thrill and murmur, a single second heart sound, and a right ventricular form of cardiac impulse. The thrill and murmur may be lower in cases of infundibular stenosis (10 per cent). In mild cases the cardiac impulse may be normal and the second heart sound clearly split.



Fig 7.24—Angiogram of a case of simple pulmonary stenosis showing dilatation of the pulmonary artery and hypoplasia of the aorta

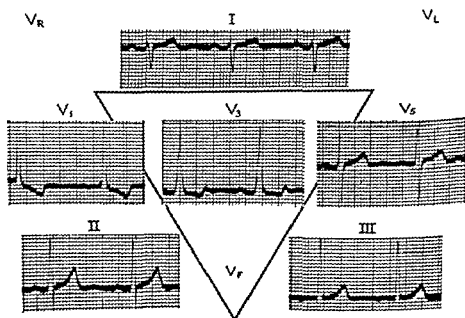


Fig 7.25—Electrocardiogram of a case of simple pulmonary stenosis showing right ventricular enlargement

Skagrams usually show dilatation of the main trunk of the pulmonary artery but not of its branches hypoplasia of the aorta and left ventricle and enlargement of the right (fig 7 24) The electrocardiogram may exhibit the usual pattern associated with marked right ventricular dominance (fig 7 25) and tall sharp P waves similar to those of chronic pulmonary heart disease but in mild cases it is normal

A high right ventricular pressure and low or normal pulmonary artery pressure may be demonstrated by means of cardiac catheterisation

Diagnosis When the chief features of a case conform to the above description the diagnosis of simple pulmonary stenosis is probable but sub aortic stenosis atrial septal defect and ventricular septal defect must be carefully excluded A common source of confusion is encountered in children with poorly developed chests who present a loud pulmonary systolic murmur and vibration The mechanism may depend upon the proximity of the anterior chest wall to the pulmonary artery in thin flat chested individuals Confusion is greater when skagrams reveal prominence of the pulmonary artery and electrocardiograms right axis deviation The final proof may rest with cardiac catheterisation

Investigation of apparently typical cases by this method may also provide proof of associated VSD or ASD when not suspected Five examples (2.5 per cent) have been encountered by the author high right ventricular pressures and relatively low pulmonary artery pressures confirming the diagnosis of pulmonary stenosis and partial admixture of arterial blood into samples obtained from appropriate chambers proving the existence of a left to right shunt Reversal of this shunt is possible in severe cases Patent foramen ovale without interauricular shunt was also discovered during cardiac catheterisation in one of the author's cases This was a girl of 14 with a mean pulmonary artery pressure of 8 mm Hg and a right ventricular pressure of 32 The pressure in the right auricle was 3 in the left 4 mm Hg If the stenosis becomes a little more severe the left auricular pressure is likely to fall and the right to rise she will then develop an interauricular shunt and become cyanosed

Prognosis Bacterial endocarditis and pulmonary tuberculosis are important risks Patients who survive these two evils may reach middle age before dying from right ventricular failure but the majority succumb when still young The average age of death in Abbott's post mortem series was 20.6 years the upper limit being 57

TREATMENT

Pulmonary valvulotomy (Brock 1948) should be considered in severe cases but carries too great a risk at present for the majority Life may be prolonged by a sedentary occupation and by limitation of effort Complications may be prevented by guarding against dental sepsis and throat infection by prophylactic chemotherapy when indicated and by annual radiological examination of the chest in order to detect early pulmonary tuberculosis at a stage when it may be successfully treated

PULMONARY STENOSIS WITH REVERSED INTERAURICULAR OR INTERVENTRICULAR SHUNT (incidence 2.5 per cent)

Relatively mild or moderately severe cases of pulmonary stenosis may have a patent foramen ovale that is functionally closed. A catheter may be passed through the valve flap into the left auricle and samples prove the absence of a shunt as described on page 242.

As life advances however pulmonary stenosis may become more severe the pressure in the right side of the heart then rises and the foramen ovale may suddenly begin to function and permit the passage of blood from right to left auricle. Such cases illustrate very well what is really meant by late central cyanosis or cyanose tardive. Patients with simple pulmonary stenosis and ASD or VSD may behave similarly.

The change is apt to occur in the teens or early twenties and is accompanied by the development of breathlessness and sometimes by syncope on effort. Cyanosis is notably variable and patients may turn almost black on exertion. Simple pulmonary stenosis with reversed interventricular shunt resembles Fallot's tetralogy but the aorta is not overriding.

In the most severe cases of pulmonary stenosis the pressure in the right auricle exceeds that in the left from birth and the foramen ovale cannot close. Under these circumstances patients have permanent central cyanosis and proportionate functional incapacity from birth.

The clinical findings in cyanosed cases differ little in other respects from severe cases without cyanosis but skiagrams may show diminished vascular markings in the lung fields (fig. 7.26a) and the heart is usually larger (fig. 7.26b). Despite the theoretical consideration that the left ventricle is the better filled chamber the cardiac impulse retains its right ventricular quality and the electrocardiogram usually shows extreme right ventricular dominance (fig. 7.27).

It will now be appreciated that there are three main functional types of simple pulmonary stenosis—acyanotic without shunt (with or without patent foramen ovale), acyanotic with direct left to right shunt (through an atrial or ventricular septal defect) and cyanotic with reversed shunt (usually interatrial via a patent foramen ovale but occasionally through an atrial or ventricular septal defect). The stenosis is valvular in 90 per cent and subvalvular in 10 per cent and occasionally there is a separate infundibular chamber. Cyanosed cases are always severe, acyanotic cases may be mild or severe.

Cardiac catheterisation revealed low mean pressures in the pulmonary artery (below 8 mm Hg) and high mean pressures in the right ventricle (over 30 mm Hg) in severe cases whether cyanosed or otherwise and normal mean pulmonary artery pressures (over 8 mm Hg) and moderately raised right ventricular pressures (under 30 mm Hg) in the mild cases.



(a) Showing dilatation of the pulmonary arc and pulmonary ischaemia



(b) Showing considerable cardiac enlargement and pulmonary ischaemia

Fig 7 2f—Pulmonary valvular stenosis with reversed interatrial shunt

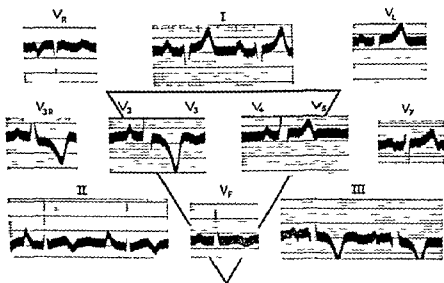


Fig 7 27—Strong right ventricular dominance due to pulmonary valvular stenosis with reversed interatrial shunt

Typical findings in a severe cyanosed case are illustrated in fig 7 28 Samples from the pulmonary veins were fully saturated with oxygen but left auricular samples were only about 70 per cent saturated, showing gross admixture of venous blood via the defect Samples from the left ventricle and femoral artery were similar and thus excluded Fallot's tetralogy

The position of the catheter when being withdrawn through a large foramen ovale or atrial septal defect is characteristic its transverse segment

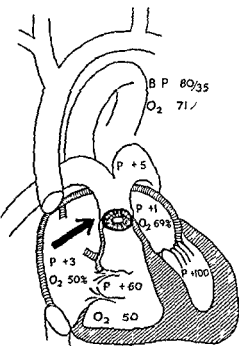


Fig 7 28—Findings on cardiac catheterization in a typical case of pulmonary valvular stenosis with reversed interatrial shunt (Pressures in cm. of saline)



Fig. 7 29—Catheter penetrating the left atrium through an atrial septal defect showing the high position of the latter (case of pulmonary stenosis plus ASD)

riding remarkably high in the centre of the heart shadow (fig 7 29) The position of the catheter when it enters the various pulmonary veins is also characteristic (fig 7 30) and is usually coiled on the right side It may be added that owing to the direction of the interatrial shunt the catheter tends to pass through the defect very easily perhaps more readily than through the tricuspid valve

Angiocardiography confirms the interatrial shunt by showing immediate filling of the left heart Owing to simultaneous opacification of both ventricles it is very difficult to obtain an unobstructed view of the outflow tract of the right ventricle and it is usually impossible to confirm the site of the stenosis by this means



(a) Right upper



(b) Right lower



(c) Left upper

Fig 730—Catheter lying in the pulmonary veins

TREATMENT

Pulmonary valvulotomy is advised in all cyanotic cases with valvular stenosis the outlook being otherwise grave. By raising the pressure in the pulmonary artery and left auricle and lowering it in the right side of the heart valvulotomy closes or reverses the interatrial shunt (depending on the anatomy of the defect in the atrial septum). If the obstruction to the right ventricle has been sufficiently relieved that chamber should be able to cope with its increased stroke volume if not it is liable to fail.

The Blalock-Taussig operation is not advised because the extra work falling on the unrelieved right ventricle consequent upon closure of the foramen ovale or reversal of the interatrial shunt usually causes early death from heart failure. Such cases appear to have shown however that increasing the pulmonary artery flow does raise the left auricular pressure.

FALLOT'S TETRALOGY

The combination of pulmonary stenosis, patent interventricular septum riding aorta and enlargement of the right ventricle is known as Fallot's tetralogy (Fallot 1888) and accounts for 75 per cent of cases of congenital heart disease with clubbing of the fingers, polycythæmia and permanent central cyanosis. The stenosis is usually subvalvular and is due to great narrowing and distortion of the right ventricular outflow tract. The pulmonary artery instead of being dilated as in simple pulmonary stenosis is remarkably small and may resemble a vein. By riding aorta is meant displacement of the root of the aorta to the right (dextroposed aorta) so that it sits astride the septum and appears to arise as much from the right as from the left ventricle. The association of these three malformations is no accident but depends upon the same embryological defect, the fault lying with arrested evolution of the bulbus cordis with incomplete torsion. A right-sided aortic arch is found in about 25 per cent of cases (Blalock 1948).

Hæmodynamics. Aortic blood is arterio-venous being composed of the full output of the left ventricle and part of that from the right. The right ventricle has a double burden for it must work against a constricted outflow tract and compete with the left ventricle against the systemic blood pressure. The situation is met by great hypertrophy of the right ventricle, the fourth constant finding in the tetralogy. The deficient pulmonary circulation is occasionally improved by extensive development of the bronchial vascular system. Polycythæmia helps to compensate for anoxæmia. Occasionally the defect is partly corrected by patency of the ductus but a continuous murmur on either side of the chest usually signifies pulmonary atresia with an extensive broncho-pulmonary anastomosis.

CLINICAL FEATURES

Fallot's tetralogy including three cases of pulmonary atresia occurred in 18 per cent of the author's series. This figure is probably influenced by slight weighting in favour of anomalies amenable to surgical repair.

Cyanosis, polycythemia and clubbing of the fingers may be absent in infants but develop in early childhood and tend to be progressive. Growth may be stunted and mental development retarded in severe cases. The chief symptom is breathlessness and to obtain maximum comfort children often adopt a characteristic squatting posture (Taussig 1947). Improve



(a) Antero posterior view

Fig 731—Skigram of a case of Fallot's tetralogy showing the characteristic



(b) Second oblique position

ment sometimes occurs owing to the development of a bronchial collateral circulation.

The pulse is apt to be small, the jugular venous pressure normal and giant a waves unusual. The cardiac impulse is tapping and no pulmonary artery pulsation can be detected. A systolic murmur is usually heard, being high in one third and in the Roger area in the remainder.

A thrill is felt in half the



Fig 732—Right-sided aortic arch in a case of Fallot's tetralogy



Fig 7 33—Angio ardiogram of a case of Fallot's tetralogy showing simultaneous opacification of the aorta and pulmonary artery i.e. a right left shunt. The stenosis appears to be subvalvular. Both pulmonary and both subclavian arteries can be seen.

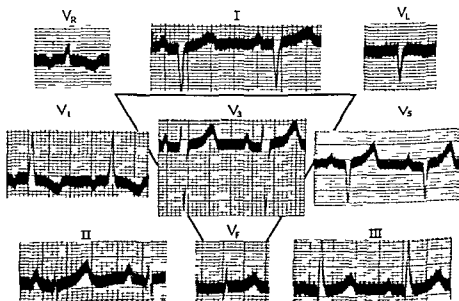


Fig 7 34—Electrocardiogram of a case of Fallot's tetralogy showing marked right ventricular dominance.

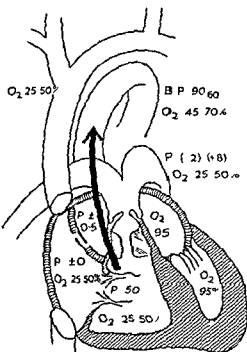


(a) Main trunk



(b) Right pulmonary artery (1st oblique view) (c) Left pulmonary artery (2nd oblique view)

Fig 7 35—Catheterization of the pulmonary artery in a case of Fallot's tetralogy



FALLOT'S TETRALOGY

Fig 7 36—Average catheter findings in Fallot's tetralogy (Pressures in cm. of saline)



Fig 7 37—Penetration of the foramen ovale and pulmonary veins in a case of Fallot's tetralogy

cases and may be high or low. The second sound at the base is single never split owing to the absence of the pulmonary element and may be quite loud.

The skiagram is usually pathognomonic and is characterised by conspicuously clear lung fields due to diminution of the pulmonary vascular markings by a notable gap between the aortic knuckle and ventricles due to hypoplasia of the pulmonary artery and by a tip tilted cardiac apex (fig 7 31). This is the *cœur en sabot* for it resembles the shape of a peasant's wooden shoe with turned up toe. The effect is produced by considerable hypertrophy of the right ventricle with displacement of the interventricular septum to the left so that the left ventricle appears as a small cap above the right ventricular apex. Occasionally, the lung fields present a reticular appear-



Fig 7 38—Site of abrupt pressure change in pulmonary valvular stenosis



Fig 7 39—Pulmonary subvalvular stenosis
the tip of the catheter is still in a low pressure
zone



(a) Catheter lying in right pulmonary artery
which can be seen clearly

(b) Left pulmonary artery [not obvious in
(a)] outlined after injecting 5 ml of 70 per
cent dioxane through the catheter

Fig 7 40—Demonstration of the anatomy of the pulmonary arteries in a case of Fallot's tetralogy

ance representing development of the bronchial circulation. In the left anterior oblique view the heart shadow may be globular, owing to the increased curvature of the right auricle and ventricle. If the aorta is right sided, the knuckle may be seen above the right auricle (fig 7 32) and the barium filled œsophagus is deflected to the patient's left. The shunt may be demonstrated by means of angiocardiology (Grishman, Steinberg and Sussman, 1941) which shows immediate filling of the aorta and great vessels and undersized pulmonary arteries (fig 7 33). The antero-posterior view is advised in order to reveal the anatomy of both subclavian arteries—a point of considerable interest to the surgeon.

The electrocardiogram shows extreme clockwise rotation or right ventricular dominance (fig 7 34), and often the tall sharp P wave of right auricular hypertrophy.

Cardiac catheterisation may be helpful in doubtful cases. The patient should receive 50 to 100 mg of heparin at the start of the procedure in order to minimise the risk of paradoxical thrombo-embolism and great care must be taken not to introduce air into the system. Rigid asepsis should also be maintained and adequate doses of penicillin should be given for forty-eight hours to prevent the possibility of subsequent cerebral abscess.

In twenty-one cases studied by the author the mean right ventricular pressure ranged between 23 and 56 mm Hg and averaged around 40. The pulmonary artery was entered in fifteen cases (fig 7 35) the pressures falling sharply to less than 8 mm Hg above the sternal angle, the range being -1 to +7 and the average +3.5 (fig 7 36). The right auricular pressure was usually close to zero (-1 to +3). A patent foramen ovale was penetrated in three cases (fig 7 37) the left auricular pressure was higher than the right on each occasion and samples from the pulmonary veins and left auricle were about 95 per cent saturated with oxygen showing that there was no right to left interatrial shunt. The similarity of samples from the venæ cavæ, right auricle, right ventricle and pulmonary artery proved the absence of left to right shunt; all these samples were grossly unsaturated (94–212 ml per L). Samples from the femoral artery were mostly between 45 and 75 per cent saturated with oxygen.

Catheterisation not only proved the diagnosis and degree of Fallot's tetralogy in these cases but showed whether the stenosis was valvular or subvalvular. In valvular stenosis the change from right ventricular to pulmonary artery pressure and pulsation was abrupt and occurred when the tip of the catheter passed beyond the level of the left atrial appendage, a horizontal line through which has been found to mark the level of the pulmonary valve in controls (fig 7 38). In subvalvular stenosis the pressure changed at a lower level and sometimes less abruptly (fig 7 39).

Cyanosis, polycythæmia, clubbing and a low arterial oxygen saturation prove the existence of a veno-arterial shunt. The size of the shunt may be calculated by subtracting the pulmonary blood flow from the tricuspid (systemic) blood flow.

Owing to the scanty pulmonary blood flow small quantities of 70 per cent diiodone (5 to 10 ml) introduced directly into the pulmonary artery via the catheter opacify the pulmonary vessels very well (fig 7 40)

The demonstration of an arm to tongue circulation time of less than six seconds (McGuire and Goldman 1937) is practically valueless the test being unreliable unnecessary and apt to thrombose precious veins

The conspicuous rise in arterial oxygen saturation that occurs when a patient with Fallot's tetralogy is placed in an oxygen tent has been confirmed by most workers and has not yet been satisfactorily explained The thesis that the function of the pulmonary epithelium is impaired cannot be maintained in view of the fully oxygenated samples of blood obtained from the pulmonary veins in cases with patent foramen ovale

Prognosis Uncomplicated relatively mild cases may reach middle life but the majority die young According to Campbell (1948) only one patient in ten with congenital cyanotic heart disease reaches the age of 24 and only one in five reaches the age of 12 Bacterial endocarditis pulmonary tuberculosis pyogenic respiratory infection and cerebral abscess from paradoxical embolism (Robbins 1945) are the most serious complications but the majority appear to die from asphyxia Congestive failure is rare

TREATMENT

Prophylactic therapy should be directed against infection and overwork Pyogenic pulmonary infection may require treatment in an oxygen tent Venesection is contraindicated for dangerous anoxæmia and syncope may result

As a result of Taussig's observation that infants with Fallot's tetralogy deteriorated when the ductus arteriosus closed and that cases complicated by persistent patent ductus fared better than those without she and Blalock devised the anastomotic operation which has proved so successful (Blalock and Taussig 1945 Blalock 1946) One or other subclavian artery is anastomosed to the homolateral branch of the pulmonary artery Better alignment is obtained as a rule with the right subclavian but the left has a longer intrathoracic course and is therefore easier to bring down If results are poor a second anastomosis may be carried out later on the opposite side

Another method of achieving the same object is to make a direct anastomosis between the aortic arch and the left pulmonary artery (Potts *et al* 1946 1948)

Selection of cases for operation should be based on the following criteria

- 1 The diagnosis should be proved
- 2 The patient should have sufficient disability (apart from cosmetic discontent)
- 3 The arterial oxygen saturation should be less than 75 per cent at rest
- 4 The mean pulmonary artery pressure should be close to zero certainly less than 10 cm of saline

- 5 Both pulmonary arteries should have been demonstrated
- 6 Both subclavian arteries should have been demonstrated
- 7 The anatomy of the aortic arch should have been demonstrated

Angiocardiography and cardiac catheterisation supply most of the required information

According to Bing (1947) the oxygen uptake per litre of ventilation does not increase during effort in Fallot's tetralogy or in other cyanosed cases of congenital heart disease that would be improved by the Taussig-Ballock operation and the test is being used to aid the selection of cases suitable for surgery

The best age at which to operate is between 3 and 10 years. The physiological results of technically successful anastomosis are good: cyanosis and clubbing may disappear, breathlessness decreases, the habit of squatting is usually given up and effort tolerance improves, the arterial oxygen saturation rises to the region of 80 per cent and the blood count returns to normal (Taussig 1948). A loud machinery murmur and coarse thrill may be detected on the homolateral side immediately after the operation in nearly all cases and are permanent. The risks of pulmonary tuberculosis and bronchopneumonia are likely to be diminished but the incidence of bacterial endocarditis should not be altered.

Ballock's total mortality rate for this operation is 17 per cent but this includes infants (mortality rate 25 per cent), cases of tricuspid atresia and other anomalies. His mortality rate for selected cases of Fallot's tetralogy is not more than 10 per cent.

If the stenosis is valvular, an alternative operation is pulmonary valvulotomy (Brock 1948). The results in successful cases are satisfactory but the risks appear to be heavier. Further technical experience however may reduce the mortality rate. Moreover certain types of subvalvular stenosis may also be amenable to direct attack.

EISENMENGER'S COMPLEX

Relatively rare cases occur in which 'riding aorta', patent interventricular septum and right ventricular enlargement are associated with a normal right ventricular conus, normal pulmonary valve and dilated pulmonary artery (Eisenmenger 1897). The riding aorta allows venous blood to escape into the systemic circulation so that slight to moderate central cyanosis occurs. The prognosis is much better than in Fallot's tetralogy, the majority of cases attaining middle life.

The physical signs include a systolic thrill and murmur maximum at the third left intercostal space near the sternal border, palpable pulsation over the pulmonary artery, accentuation of the second or pulmonary element of a normally split second sound at the base and occasionally a pulmonary or aortic diastolic murmur.

Right ventricular enlargement and dilatation of the pulmonary artery

may be demonstrated by means of electrocardiography and X rays (fig 7 41) The arterial oxygen saturation is reduced usually to 70 to 80 per cent right ventricular and pulmonary artery pressures are raised right ventricular and pulmonary artery samples are similar to samples from the right auricle and superior vena cava and angiocardiology (fig 7 42) shows simultaneous opacification of the aorta and pulmonary artery



Fig 7 41—Eisenmenger's complex

(By courtesy of Dr Maurice Campbell)



Fig 7 42—Angiogram of a case of Eisenmenger's complex showing simultaneous opacification of the aorta and dilated pulmonary artery

(Sussman and Grishman 1947) In two cases investigated by the author and proved with the aid of angiocardiology the shunt was only from right to left samples from the pulmonary artery being the same as those from the right side of the heart The arterial oxygen saturation was 72 per cent in one case and 71 per cent in the other and the pulmonary blood flow was considerably reduced in both Unlike Fallot's tetralogy however the mean pressure in the pulmonary artery was high

In differential diagnosis it is well to remember that the majority of cases that appear at first to be examples of the Eisenmenger complex turn out otherwise The following should be considered

- 1 *Atrial septal defect with late reversal of the shunt* Wide splitting of the second heart sound right bundle branch left auricular blood samples containing a proportion of venous blood and angiocardigrams showing immediate filling of the left auricle distinguish it

- 2 *Pulmonary hypertension with late shunting through a patent foramen*

orale This is distinguished by catheter samples from the left auricle and by angiocardigrams as described for (1) above

3 *Idiopathic pulmonary hypertension with late reduction in arterial oxygen saturation* There is usually no pulmonary systolic murmur, pulmonary venous blood samples are improperly saturated and are similar to samples from the left auricle left ventricle and femoral artery, angiocardigrams show no shunt

4 *Anoxic pulmonary heart disease* This does not cause confusion when there is obvious advanced emphysema, but it may give rise to difficulty when the degree of emphysema seems insufficient to cause cyanosis The absence of an intracardiac shunt must then be demonstrated

5 *Transposition of the great vessels with patent septa* Cyanosis is usually earlier in onset Samples of blood from the pulmonary artery contain more oxygen than samples from the aorta or peripheral arteries and samples from the right auricle are more saturated than those from the venæ cavae Angiocardigrams may be inconclusive Skiagrams show pulmonary plethora

6 *Ventricular septal defect or patent ductus with reversal of the shunt due to pulmonary hypertension* Such cases are clinically indistinguishable from the Eisenmenger complex

TREATMENT

Blalock's operation is contraindicated because an artificial ductus cannot function well when the pulmonary blood pressure is raised nor is the arterial oxygen saturation low enough to warrant it

OTHER ANOMALIES CAUSING PERMANENT CYANOSIS

The remaining malformations listed in the classification on page 203 are rarely compatible with more than a few months of life unless associated with other anomalies that have a favourable influence on hæmodynamics

Transposition of the great vessels the pulmonary artery arising from the left ventricle and the aorta from the right results in two independent circulations Life depends principally upon an atrial septal defect whereby oxygenated blood from the left auricle passes into the right side of the heart and so into the aorta and upon a patent interventricular septum through which venous blood in the right ventricle may be pumped into the lungs

The chief presenting features are central cyanosis associated with a split second heart sound dilatation of the pulmonary artery and engorgement of the pulmonary vascular shadows (fig 7 43) The electrocardiogram reveals gross right ventricular dominance or right bundle branch block Cardiac catheterisation shows that samples from the pulmonary artery (entered from the right ventricle via the ventricular septal defect fig 7 44) contain more oxygen than samples from the aorta (also entered from the right ventricle fig 7 45) or femoral artery a state of affairs that can occur in no other

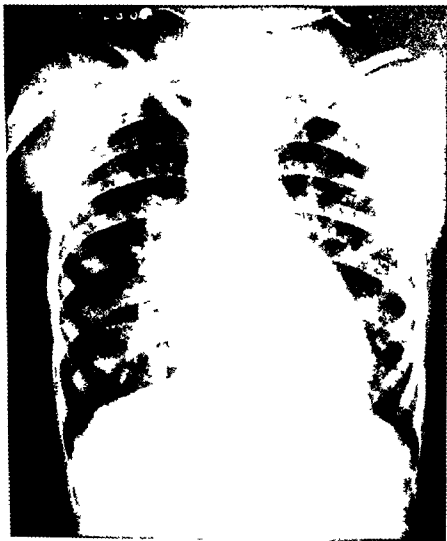


Fig 7 43—Skiagram of a case of transposition of the great vessels associated with a
and ventricular septal defects



Fig 7.44—Skiagram of a case of transposition of the great vessels showing penetration of the pulmonary artery from the right ventricle via the ventricular septal defect



Fig 7 45—Skysagram of a case of transposition of the great vessels showing a catheter penetrating the aorta directly from the right ventricle

condition Aortic samples are similar to those from the right ventricle and auricle and contain considerably more oxygen than samples from the superior vena cava by virtue of the interatrial shunt In a case investigated by the author samples from the aorta and right heart were 82 per cent saturated from the superior vena cava 60 per cent and from the pulmonary artery 89 per cent The right ventricular systolic pressure equals the aortic pressure The pulmonary blood flow is greatly increased (9 L/min in the case cited) the aortic decreased (3 L/min) Angiocardiography shows simultaneous filling of the aorta and pulmonary artery from the right ventricle A second case investigated showed no ASD but greater pulmonary plethora The catheter was again manipulated into both aorta and pulmonary artery

Persistent truncus arteriosus is due to failure of development of the aorto pulmonary septum so that a single large vessel arises from both ventricles Blood is carried to the lungs either by pulmonary arteries arising from the common trunk or by way of the bronchial arteries Cyanosis is minimal when the pulmonary arteries are present extreme when the pulmonary blood flow depends on the bronchial arteries Radiologically the aortic knuckle and vascular pedicle are unduly prominent and there is a conspicuous bay between the knuckle and the left border of the heart Both ventricles are grossly enlarged

A harsh systolic murmur and thrill are usually present at the base and the second heart sound is loud and single The electrocardiogram shows right ventricular dominance or right bundle branch block Angiocardiography helps to establish the diagnosis especially when the pulmonary arteries are absent but appearances closely resemble pulmonary atresia

A Blalock Taussig operation may benefit those cases in which the blood flow to the lung is reduced that is those cases with severe cyanosis in which the bronchial arteries supply the lungs

Tricuspid atresia with a functionless right ventricle and atrial septal defect is characterised by intense permanent cyanosis due to right to left interatrial shunt by gross left ventricular enlargement and by electrocardiographic evidence of considerable left ventricular preponderance The blood flow to the lungs is greatly reduced and depends upon a patent interventricular septum patent ductus or upon the bronchial collaterals The latter may link up directly with the larger pulmonary arteries and cause a widespread bilateral machinery murmur The diagnosis may be confirmed by means of angiocardiography (fig 746) and cardiac catheterisation and the condition may be greatly improved by the Blalock Taussig operation



(a)



(b)



(c)



(d)

Fig 746 —Angiocardiogram in case of tricuspid atresia (a) Showing dye passing directly into left auricle (1 second) (b) Showing early filling of the left ventricle and aorta (2 seconds) note bronchial collaterals in right upper zone (c) Second oblique position the left side of the heart and the aorta are filled in 2 seconds (d) At 3 seconds a small left pulmonary artery is becoming visible

REFERENCES

- Abbott M E (1948) Coarctation of the aorta of the adult type II Statistical and historical retrospect of 200 recorded cases with autopsy of stenosis or obliteration of the descending arch *Amer Heart J* 3 39. — (1934) Congenital heart disease Nelson loose leaf living Medicine 4 207
- Addarii F Martini L Mahaim I and Winston M (1946) Anatomical and clinical data in a case of irreducible cardiac insufficiency of uncertain etiology occurring in a young man New investigations on incomplete bilateral block *Cardiologia* 11 36
- Baldwin E de F Moore L V and Noble R P (1946) The demonstration of ventricular septal defect by means of right heart catheterisation *Amer Heart J* 32 152
- Bedford D E Papp C and Parkinson J (1941) Atrial septal defect *Brit Heart J* 3 37
- Benn J (1947) The prognosis of patent ductus arteriosus *Ibid* 9 53
- Bing R J Vandam L D and Gray F D (1947) Physiological studies in congenital heart disease I Procedures II Results of pre operative studies in patients with tetralogy of Fallot III Results obtained in five cases of Eisenmenger's complex *Bull Johns Hopk Hosp* 80 107 121 3-3
- Blalock A (1946) Physiopathology and surgical treatment of congenital cardiovascular defects *Bull New York Acad Med* 22 57 — (1948) Surgical treatment of pulmonary stenosis *Brit Heart J* 10 68 — Taussig H B (1945) The surgical treatment of malformations of the heart in which there is pulmonary stenosis or pulmonary atresia *J Amer med Ass* 128 189
- Bland E F White P D and Garland J (1933) Congenital anomalies of the coronary arteries report of an unusual case associated with cardiac hypertrophy *Amer Heart J* 8 787
- Bohn H (1938) Ein wichtiges diagnostisches Phänomen zur Erkennung des offenen ductus art Botalli *Klin Wchschr* 17 907
- Bonnet L M (1903) Sur la lesion dite Sténose congénitale de l'aorte *Rev Med* 23 108
- Bramwell C (1947) Coarctation of the aorta II Clinical features *Brit Heart J* 9 100
- Brock R C (1948) Pulmonary valvulotomy for the relief of congenital pulmonary stenosis *Brit med J* 1 1121
- Broden B Hanson H E and Karnell J (1948) Thoracic aortography preliminary report *Acta Radiol* 29 181
- Brown G E Clagett O T Burchell H B and Wood E H (1948) Pre operative and postoperative studies of intraradial and intrafemoral pressures in patients with coarctation of the aorta *Proc Mayo Clin* 23 52
- Brown J W (1939) Congenital heart disease London
- Campbell M (1948) The Blalock Taussig operation for morbus caeruleus *Guy's Hosp Rep* 97 1
- Cassels D E and Morse M (1947) Blood volume in congenital heart disease *J Pediat* 31 485
- Crafoord C (1948) Coarctation of the aorta *Brit Heart J* 10 71 — (1948) Patent ductus arteriosus *Ibid* 10 74 — Nylin G (1945) Congenital coarctation of the aorta and its surgical treatment *J thor Surg* 14 347
- Donovan M S Neuhauser E B D and Sosman M C (1943) The Roentgen signs of patent ductus arteriosus A summary of 50 surgically verified cases *Amer J Roentgen* 50 293
- East T (1932) Coarctation of the aorta *Proc Roy Soc Med* 25 796
- Eisenmenger V (1897) Die angeborenen Defecte der Hammerscher wand des Herzens *Ztschr f klin Med* 3. Supplementz Heft 1
- Evans W (1933) Congenital stenosis (Coarctation) atresia and interruption of aortic arch study of 28 cases *Quart J Med* 2 1 — (1947) Familial cardiomegaly *Brit Heart J* 2 309

Fallot A (1888) Contribution a l'anatomie pathologique de la maladie bleue (cyanose cardiaque) *Marseille Med* 25 77

Gibson G A (1900) Clinical lectures on circulatory affections Lecture 1 persistence of the arterial duct and its diagnosis *Edin med J* 8 1

Glynn I E (1940) Medial defects in circle of Willis and their relation to aneurysm formation *J Path Bact* 51 213

— and Reinhold J D L (1949) The relationship of idiopathic cardiac hypertrophy to foetal endocarditis (in the Press)

Grishman A Steinberg M F and Susysman M L (1941) Contrast roentgen visualisation of coarctation of the aorta *Amer Heart J* 21 365 — —

— (1941) Tetralogy of Fallot contrast visualisation of the heart and great vessels *Radiology* 37 178

Gross R E (1947) Complete division for the patent ductus arteriosus *J thor Surg* 16 314 — (1949) Surgical treatment for coarctation of the aorta Experience from 60 cases *J Amer med Ass* 139 285 — Hubbard

J P (1939) Surgical ligation of a patent ductus arteriosus *Ibid* 112 7-9

Howarth S McMichael J and Sharpey Schafer E P (1947) Cardiac catheterisation in cases of patent interauricular septum primary pulmonary hypertension Fallot's tetralogy and pulmonary stenosis *Brit Heart J* 9 292

Ingham D W and Williams F A (1938) Congenital transposition of the great arterial trunks *Amer Heart J* 15 482

Jones J C (1947) Complications of the surgery of patent ductus arteriosus *J thor Surg* 16 305

Keith A (1909) The Hunterian lectures on malformations of the heart *Lancet* ii 359

King J T (1937) The blood pressure in stenosis at the isthmus of the aorta *Ann intern Med* 10 1802

Kugel M A (1939) Enlargement of the heart in infants and young children *Amer Heart J* 17 60.

Laubry C Routier D and de Balsac R (1941) Grosse pulmonaire Petite aorte Affection congenitale *Bull et mem Soc m d d hop de Paris* 56 847

Love W S and Holms J H (1939) Coarctation of the aorta with a associated stenosis of the right subclavian artery *Amer Heart J* 17 6 8

Lutembacher R (1916) De la stenose mitrale avec communication inter auriculaire *Arch d mal d Cœur* 9 237

McGinn S and White P D (1933) Interauricular septal defect a sociated with mitral stenosis *Amer Heart J* 9 1

McGuire J and Goldman F (1937) Apparent increased velocity of blood flow in cases of congenital heart disease with septal defect having right to left shunts *Ibid* 14 230

Muir D C and Brown J W (1934) Patent interventricular septum (maladie de Roger) *Arch Dis Child* 9 27

Murray G (1948) Closure of defects in cardiac septa *Ann Surg* 128 843

Perry C B (1931) Congenital heart disease as seen in elementary school children *Bristol med chir J* 48 41

Potts W J and Gibson S (1948) Aortic pulmonary anastomosis in congenital pulmonary stenosis *J Amer med Ass* 137 343 — Smith S and

Gibson S (1946) Anastomosis of the aorta to a pulmonary artery Certain types in congenital heart disease *Ibid* 132 6-7

Railsbach O C and Dock W (1929) Erosion of ribs due to stenosis of isthmus (coarctation) of aorta *Radiol* 12 58

Reifenstein G H Levine S A and Gross R E (1947) Coarctation of the aorta A review of 104 autopsied cases of the adult type 2 years of age or older *Amer Heart J* 33 146

Robbins S L (1945) Brain abscess associated with congenital heart disease *Arch intern Med* 75 279

Roesler H (1928) Beiträge zur Lehre von den angeborenen Herzfehlern IV Untersuchungen an zwei Fällen von Isthmus stenose der Aorta *Wien Arch inn Med* 15 521 — (1934) Interatrial septal defect *Arch intern Med* 54 339

Roger H (1879) Recherches cliniques sur la communication congenital des deux cœurs par inoclusion du septum interventriculaire *Bull Acad Med d Paris* 8 1074

Rytand D A (1938) The renal factor in arterial hypertension with coarctation of the aorta *J clin Invest* 17 391

Shapiro M J (1949) Clinical studies on twenty one cases of coarctation of the aorta *Amer Heart J* 37 1045 — and Keys A (1943) Patency of ductus arteriosus in adults *Ibid* 25 158

Steinberg M F Grishman A and Sussman M L (1943) Angiocardiography in congenital heart disease III Patent ductus arteriosus *Amer J Roentgenol* 50 306

Sussman M L and Grishman A (1947) A study of angiocardiography and angiography *Advances in internal Medicine* New York 2 102

Swan C *et al* (1943) Congenital defects in infants following infectious diseases during pregnancy with special reference to relationship between German measles and cataract deaf mutism heart disease and microcephaly and to period of pregnancy in which occurrence of rubella is followed by congenital abnormalities *Med J Australia* 2 201

Taussig H B (1948) The surgery of congenital heart disease Diagnosis and treatment of the cyanotic group *Brit Heart J* 10 65 — (1947) Congenital malformations of the heart New York

Touroff A S W and Vesell H (1940) Subacute streptococcus viridans endarteritis complicating patent ductus arteriosus *J Amer med Ass* 115 170

Vesell H and Kross I (1946) Patent ductus arteriosus with subacute bacterial endarteritis diagnosis and indications for operation *Arch intern Med* 77 659

Weber F (1918) Can the clinical manifestations of congenital heart disease disappear with the general growth and development of the patient? *Brit J child Dis* 15 113

Wilson M G and Iubaszcz R (1942) Prognosis for children with congenital anomalies of the heart and central vessels *J Pediat* 21 23

Wood P H (1950) Congenital Heart Disease *BMJ* (St Cyres Lecture in the press)

CHAPTER VIII

RHEUMATIC FEVER AND ACTIVE RHEUMATIC CARDITIS

RHEUMATIC fever is a particular form of polyarthritis following streptococcal infection its hall marks are pancarditis chorea subcutaneous nodules and erythema marginatum It may be acute subacute or chronic

INCIDENCE

According to the 1927 report of the Child Life Committee of the Medical Research Council Social Conditions and Acute Rheumatism 10 to 15 per cent of all children at 12 years of age in England are affected by rheumatism Of 22 800 children under 15 years of age card indexed by the London County Council 2.6 per cent had had rheumatic fever (Bach *et al* 1939) The crude annual death rate from rheumatic fever declined from 67 per million persons in 1901 to 22 per million in 1937 (Glover 1939) During 1937 according to Glover rheumatic fever accounted for 2.3 per cent of all deaths in children between the ages of 5 and 9 years

The disease is rare in infancy and in old age and is most common in childhood and adolescence attacking the poor rather than the rich and having an incidence climatically and geographically parallel to streptococcal tonsillitis (Coburn 1931) The peak incidence is in children between the ages of 8 and 12 particularly during the months of October November and of January February Apart from arachnodactyly there is no evidence that a particular physical type is predisposed to rheumatic fever (Hill and Allan 1929) but hereditary predisposition is now accepted (Wilson 1940)

THE NATURE OF THE RHEUMATIC STATE

There is no evidence as yet that rheumatic fever is caused directly by any infective agent Cultures from blood joint fluid pericardial or pleural effusions and from affected tissues are bacteriologically sterile and filtrates from similar samples are incapable of transmitting the disease when inoculated into animal or man There is still perhaps a remote possibility that a virus is responsible but the known facts are against it

On the other hand the evidence that rheumatic fever is intimately related to streptococcal infection is beyond dispute The relationship was first propounded by Poynton and Paine in 1900 They isolated a diplococcus from blood and other cultures and produced polyarthritis and carditis by injecting it into animals but the lesions were shown later to be infective not rheumatic However they confirmed the observation of Haig Brown

1) The rheumatic fever was nearly always preceded by streptococcal infection, the latent interval being 10 to 20 days (Parsons and Parry, 1931). The most convincing proof of this was later given by Sikes *et al.* (1931). The remarkable organism is always a haemolytic streptococcus (Group A). As previously noted, the incidence of rheumatic fever follows closely the geographical, social and seasonal incidence of streptococcal infection (Coburn 1931) and small epidemics of rheumatic fever in closed communities always follow epidemics of streptococcal sore throat (Glover 1931). Culpable streptococci belong serologically to group A and may produce a powerful erythrogenic toxin and haemolysins—in fact they are of the same kind as strains (Coburn and Pauli 1935). Serum from the subjects of rheumatic fever agglutinates these strains in high titre and anti-streptococcal haemolysins (antibodies excited by the antigenic properties of streptococcal haemolysins) have been found in high titre in the early stages of practically all cases of active rheumatic fever whether or not a history of streptococcal infection is obtained (Todd 1932). Most of these observations have been confirmed independently by other workers, notably Griffith (1935), Sheldon (1931) and Bradley (1932).

It is now generally believed that rheumatic fever is an abnormal tissue reaction to the products of haemolytic streptococcal infection in a sensitised individual. A number of other observations supports this hypothesis. Thus allergic polyarthritis with a latent interval of 8 to 9 days may follow the injection of foreign serum, polyarthritis may similarly follow gonococcal, dysenteric and other bacterial infections in individuals sensitised by previous attacks, associated skin lesions in rheumatic fever such as erythema multiforme strongly suggest allergy. Finally the experimental work of Rich and Gregory (1943, 1944) who succeeded in producing carditis of the rheumatic type in rabbits by injecting horse serum and of Cavellu (1947), who was equally successful in rats which were injected with an antigen consisting of killed streptococci and heart or connective tissue emulsion provides convincing evidence of the existence of an allergic form of carditis which may be related to the streptococcus and which at least resembles that seen in rheumatic fever.

The relationship between rheumatic fever and rheumatoid arthritis is still uncertain. Serum from patients with rheumatoid arthritis commonly agglutinates all strains of haemolytic streptococci in high dilution (Dawson *et al.* 1932) but does not as a rule contain the high titre anti-haemolysins characteristic of rheumatic fever (Stuart Harris 1935), nevertheless the anti-streptolysin titre is much higher than in normal controls (Goldie and Griffiths 1936). Whether there is any essential difference between the pathology of the affected joints and in the structure of subcutaneous nodules in the two conditions other than those which might be due to the age of the patient or to the chronicity of the lesion is still a matter of controversy (Goldie 1938). The incidence of a rheumatic type of cardiac lesion in rheumatoid arthritis is difficult to assess from the literature but appears to

range between 3 and 30 per cent in clinical studies and between 25 and 66 per cent in post mortem studies (Rogen 1947). These figures are probably too high. The subject is well reviewed by Bywaters (1950).

PATHOLOGY

In a fulminating attack which ends fatally within two or three weeks tissue microscopy reveals only non specific lesions consisting of œdema fragmentation of collagen leucocytic infiltration hyperæmia and capillary hæmorrhage (Coburn 1933). Similar lesions may occur in most acute infections toxæmias and allergic states and represent the Arthus phenomenon (Werner 1938). Many tissues are so affected particularly the synovial membranes of the larger joints the pericardium myocardium and endocardium the pleura and lung. Petechiæ may be seen clinically in the skin (purpura rheumatica) or in the ocular fundi and at autopsy they are often most obvious in the pericardium and pleura. Inflammatory œdema of soft tissues may be seen clinically independent of arthritis. Effusion into the large joints and sometimes into the pericardial or pleural cavities is characteristic and is the best example of the exudative type of lesion.

The specific rheumatic lesion however is proliferative and occurs rather later. It is characterised by the Aschoff node (Aschoff 1904). This is a small collection of large often multinucleated reticulo endothelial cells mixed with lymphocytes and plasma cells surrounding a necrotic collagenous centre. There is also fibroblastic proliferation. Whilst it is in no sense perivascular it lies in close relationship to a vessel. This lesion is particularly well seen in the myocardium. Another example of the proliferative lesion is the subcutaneous nodule which may be regarded as an aggregation of Aschoff nodes with fibroblastic tissue predominating. Macroscopic nodules may be seen occasionally on the surface of the heart (fig. 8.01).

Occasionally vascular lesions are found in the viscera which show all the features of panarteritis. Involvement of the cerebral pulmonary coronary and mesenteric arteries has been described (Ritchie 1939). Secondary thrombosis may occur but is uncommon. Later the media may become calcified.

Rheumatic inflammation of the heart valves is a true valvulitis the baneful agent entering the valve through the minute vessels which supply it (Shaw 1929). There has been considerable disagreement concerning the vascularity of normal and diseased heart valves. Langer (1937) first demonstrated the dependence of valvular blood vessels upon the presence of muscle. He showed that vessels and muscle fibres reached the free edge of the valve in the foetus and new born child but soon regressed. Also that diseased valves were frequently vascularised whereas normal adult valves were not. Gross and Kugel (1921 1925-26 1927-28 1931) who studied the coronary circulation in detail by means of radiography after injecting a barium sulphate gel confirmed Langer's observations. They also found

that in the foetus the pulmonary valve was the one best provided with muscle and blood vessels whereas in children it was the aortic cusp of the mitral valve. The fact that endocarditis *in utero* usually affected the pulmonary valve whereas in children it usually affected the mitral valve thus appeared to be understood. The decreasing incidence of valvulitis as age advanced was similarly explained. More recent work based on the injection of Indian ink instead of barium gel however has thrown doubt on these

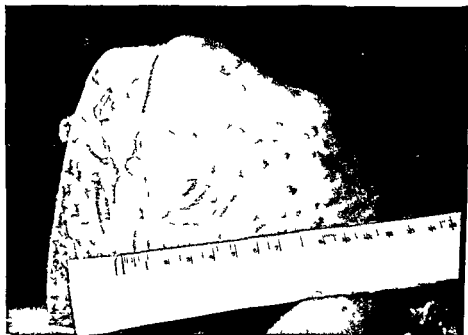


Fig. 801.—Macroscopic nodules on the surface of the heart

conclusions. Wearn *et al* (1936) for instance found capillaries in the valves of 84 per cent of seventy four normal hearts and were unable to correlate the relative incidence of endocarditis of a given valve with the frequency with which it contained blood vessels. Thus the mitral valve was vascularised in 66 per cent the tricuspid in 64 per cent the pulmonary in 28 per cent and the aortic in 16 per cent.

In the acute stage of rheumatic inflammation the valve is œdematous and soon shows signs of damage just proximal to its free edge where the cusps come into apposition i.e. at the site of maximum natural trauma. Small thrombi form on the valve at this site giving rise to a ridge or to a row of small pink nodules. As the inflammation subsides secondary sclerosis follows and results in thickening and shortening of the chordæ tendineæ in thickening and distortion of the cusps and especially in fibrosis and constriction of the mitral ring at the base of the valve. According to Carey Coombs (1924) mitral stenosis usually takes 2 to 8 years to develop. If the

mitral ring is spared and the brunt of the attack falls upon the chordæ tendineæ and the periphery of the cusps mitral incompetence may occur without stenosis and may become extreme if ring dilatation follows owing to dilatation of the left ventricle Rheumatic aortic valvulitis results in thickening and distortion of the cusps with fusion of the commissures secondary calcification and stenosis are common Tricuspid valvulitis with late stenosis or incompetence is by no means rare

CLINICAL FEATURES

In childhood the heart often bears the brunt of the attack and indeed the joints may escape entirely Once the heart has been involved however carditis or valvulitis should be assumed in all subsequent attacks the increased vascularity of a valve which has been subjected to rheumatic inflammation may partly explain this tendency to recurrence If the first attack occurs over the age of 21 carditis is unlikely and becomes progressively rare with advancing years although it may still occur even in old age Polyarthritis on the other hand becomes increasingly common

The diagnosis of rheumatic carditis is based on three major issues upon signs of some inflammatory process upon evidence that this process is rheumatic and upon proof of cardiac involvement

SIGNS OF SOME INFLAMMATORY PROCESS

These are fever leucocytosis and elevation of the erythrocyte sedimentation rate Fever may be of any degree but is usually moderate or high initially in children and moderate or low grade in adults it is irregular in type and inclined to relapse it may last only a few days or it may continue for months The temperature is normal in subacute rheumatism and may be normal when polyarthritis is still active in acute attacks Leucocytosis is slight to moderate in the early phase of acute rheumatic fever figures of 10 000 to 15 000 white cells per c mm being the rule The differential count may show a slight relative increase of polymorphs but is often normal In subacute rheumatism the total count is commonly between 7 000 and 10 000 per c mm The sedimentation rate is by far the most valuable evidence of some active inflammatory process and is often remarkably high when there are no other signs Weekly readings have proved a reliable index of the course of the disease and of the degree of activity

It will be appreciated that these three features are non specific they point to some inflammatory process but they do not determine its nature Secondary anaemia and loss of weight or failure to gain weight may be regarded in a similar light Undue tachycardia presents a more controversial problem whilst non specific for the most part and often due to anxiety it may undoubtedly result from rheumatic carditis and under certain circumstances may provide good evidence of it

EVIDENCE THAT THE INFLAMMATORY PROCESS IS RHEUMATIC

1 *Polyarthrititis* Non suppurative polyarthrititis with sterile effusions into the large joints is characteristic. Involved joints may be painful, swollen, hot, flushed, and tender, on the other hand slight effusion into a knee joint may be detected when there are no other signs or symptoms or the patient may complain of joint pains when there are no signs, as in subacute rheumatism. The older the patient the more often are the small joints affected and it becomes increasingly difficult to distinguish rheumatic fever from rheumatoid arthritis. Pains and effusions tend to fit from joint to joint, one recovering as another is involved but not necessarily. Occasionally, one knee or other large joint alone is inflamed especially if previously injured and may remain so for weeks or even for months but minimal pains elsewhere may suggest its true nature. Other forms of what is thought to be allergic polyarthrititis such as the dysenteric variety may be indistinguishable except on other grounds. For example, dysenteric polyarthrititis is proclaimed by associated conjunctivitis and urethritis and by its relation to dysentery.

In subacute rheumatism recurrent joint pains occur without effusion and usually without fever or leucocytosis but the sedimentation rate is raised. Growing pains confined to the hips, knees or ankles mean subacute rheumatism, growing pains described in the muscles, ligaments or tendons probably do not (Hawksley 1939).

2 *Relationship to streptococcal infection* The diagnosis is favoured if the symptoms follow a streptococcal sore throat or some other streptococcal infection including scarlet fever. There is a latent interval of 1 to 3 weeks usually 10 to 14 days. The significance of this relationship cannot be overstressed. It appears to be fundamentally the same as the relationship between dysenteric polyarthrititis and acute bacillary dysentery or between gonococcal polyarthrititis and gonorrhœa. Opportunity to study the dysenteric form was afforded by its frequency amongst the troops in North Africa and Italy in the second world war. It was characterised by acute polyarthrititis involving the large joints, by resistance to salicylates, by prolonged activity averaging about three months and by associated conjunctivitis and urethritis. Joint effusions were sterile and cultures from the conjunctiva and urethra yielded no pathogenic organisms. The provocative attack of dysentery was often abortive or very mild and the latent period 10 to 14 days. A previous attack of dysentery was invariable and was usually untreated. The evidence suggested that a fairly high degree of immunity was necessary for the development of the syndrome. Gonococcal polyarthrititis was equally common and behaved similarly except that tenosynovitis replaced conjunctivitis and urethritis was primary. The facts suggest that rheumatic fever is streptococcal polyarthrititis and bears the same relationship to the streptococcus as does dysenteric polyarthrititis to the dysentery bacilli and gonococcal polyarthrititis to the gonococcus but instead of con

conjunctivitis urethritis or tenosynovitis there may be carditis chorea or marginate erythema

If there is no history of recent sore throat or other streptococcal infection evidence of such may be afforded by an anti streptolysin titre in the region of 200 Todd units. High titres do not prove that an illness is rheumatic fever only that there has been recent hæmolytic streptococcal infection. Similar proof may be obtained by finding that the patient's serum agglutinates an emulsion of hæmolytic streptococci at a titre of 1:200. It is highly improbable that any case of acute rheumatic fever whether it be the first attack or a recurrence will not show such serological changes.

That continued hæmolytic streptococcal infection is not responsible for the disease may be proved by the lack of improvement after treatment with penicillin.

3 *Response to salicylates* Joint pains and effusions in rheumatic fever commonly respond dramatically to sodium salicylate in initial doses of 15 to 20 grains (1 to 1.5 G) three hourly. Aspirin is also effective. Other forms of polyarthritis (except perhaps that due to serum sickness) are not improved. Only the exudative lesion and the associated fever respond; no effect is observed on proliferative lesions. Sodium salicylate is often used as a diagnostic test but although a good one it is not infallible.

4 *Chorea* Rheumatic or Sydenham's chorea is mysterious in several ways. First it has a solitary nature preferring to occur alone rather than in the company of other rheumatic manifestations. Secondly it does not affect the sedimentation rate. Thirdly there is no specific rheumatic pathology in the brain (Shaw 1929). Nevertheless it is certainly part of the rheumatic state. About 20 per cent of patients with chorea alone develop rheumatic heart disease; about 50 per cent develop other rheumatic manifestations with or without carditis (Sutton and Dodge 1938) and most of the remainder have a familial link. Clinical features include spontaneous involuntary inco-ordinated movements, muscular weakness and alteration of tendon jerks, emotional instability and some disturbance of higher cortical function. Occasionally it is more or less confined to one side of the body. Movements disappear during sleep.

The diagnosis must be made from common tics and from hysteria in general. Reliance should be placed on the quality of the movements. They are quick, complicated, elaborate, irregular and varied. The same movement is rarely repeated exactly. The hands writhe and twist, the patient trying to stop them or attempting to conceal them by some volitional act. She often drops things she is holding, especially crockery, or she is clumsy in other ways. Facial grimaces are odd and varied, unlike the repeated twitch of a tic. After protruding the tongue for inspection she withdraws it like a lizard, snapping the jaws over it. When the hands are held out the wrist is flexed and the fingers hyper-extended. The knee jerk may be sustained, the leg being held up at the height of its extension for an appreciable interval before relaxation occurs.



(a)



(b)

Fig 80 —Erythema marginatum

Hysterical movements are more jerky and show constant repetition. Experience and familiarity with both conditions usually makes their distinction easy. The involuntary athetotic movements of encephalitis and Wilson's disease may be more confusing.

5. *Skin lesions.* Petechiae may occur in the skin or in the fundi in fulminating cases but are in no way specific. Urticaria, erythema nodosum



Fig. 803—Erythema marginatum

and erythema multiforme are sometimes seen but they too are not specific. They are probably allergic skin reactions and when associated with rheumatic fever may depend upon skin sensitisation to the streptococcus or to its toxins. Urticaria may be due to a host of antigens, erythema nodosum to the tubercle bacillus, the meningococcus or other organisms, erythema multiforme is perhaps more closely related to the streptococcus.

Erythema marginatum (Barlow and Warner 1881) a variety of erythema multiforme is especially important because it is peculiar to the rheumatic state (Cheadle 1889). It appears in rings, crescents, ovals or in irregular forms characterised by a thin red margin outlining a patch of apparently normal skin (figs 802a and b). It is distributed chiefly over the trunk and proximal part of the limbs. There may be two or three lesions or dozens of them. Sometimes the rash is at first composed of irregular erythematous macules (fig 803) but the centres soon clear leaving spreading red margins (Perry 1937). Erythema marginatum may be fleeting or remarkably persistent; as a rule it is recurrent and may reappear from time to time long after other manifestations of active rheumatism have subsided.



(a) On the knuckles



(b) On the back of the head

Fig 8 04—Subcutaneous rheumatic nodules

Subcutaneous nodules are good examples of proliferative rheumatic lesions. Like erythema marginatum they were first properly studied by Barlow and Warner (1881). Varying in size from something so small as to escape clinical perception to the dimensions of a Barcelona nut they are usually attached to tendon sheaths to the superficial surface of joint capsules or to other fascia so that the skin rides over them freely. Sometimes they are partially attached to the skin and more or less move with it. They are best seen on the knuckles (fig 8 04a) on the back of the head (fig 8 04b) on the elbows or on the knees. In children they are practically diagnostic of rheumatic fever but similar though usually larger and more persistent nodules may occur in Still's disease and in adult rheumatoid arthritis (Hawthorne 1900). It is doubtful whether there is any fundamental difference between these nodules (Keil 1938).

6 Pulmonary signs Because of its non specific clinical features pleurisy rarely provides evidence of rheumatic fever but it is not uncommon. Paul (1928) gave its incidence as 10 per cent. It may be dry or it may give rise to a sterile straw coloured effusion. Response to salicylates is in different

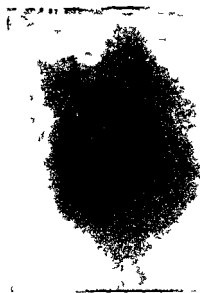


Fig 8 05—Skiagraph showing rheumatic pneumonia in a girl

Rheumatic pneumonia is rare occurring in only 1 to 2 per cent of active cases. Symptoms are not spectacular. There is no chill, breathing is not embarrassed, the respiratory rate is but little elevated and fever is not necessarily higher than before. Cough may be noted but is rarely troublesome. The sputum is scanty and tenacious, occasionally it is streaked with blood. Physical signs include dullness to percussion, bronchial breathing and crepitations appearing first here then there. The transient and migratory nature of these signs is characteristic. Serial skiagraphs confirm the presence of patchy wandering consolidation or may show a variable broncho-pneumonic pattern (fig 8 05). The white blood count is little altered. Rheumatic pneumonia is not influenced by penicillin, sulphonamides or salicylates; fortunately it does not often appear to alter the course of the major illness.

Most cases studied at autopsy have been unusually severe and consolidation has been extensive and mostly lobar in distribution. The affected parts

are bulky have a peculiar succulent gelatinous appearance (Hadfield 1938) and feel like indiarubber in colour they are a homogenous rich purplish red (Naish 1928 Liman and Gouley 1928) and later may be buff Microscopically the predominant finding is an extensive fibrinous exudate infiltrated with mononuclear and multinucleated cells Polymorphs and lymphocytes are scanty The cellular exudate is partly interstitial but also lines the alveolar ducts and may fill the alveoli (Hadfield 1938) There is associated hyperæmia and œdema Secondary fibroblastic reaction develops later and when interstitial may be responsible for pulmonary hypertension (Gouley 1938) Similar lesions have been produced experimentally by Rich and Gregory (1943)

Simple collapse of either lower lobe in the course of rheumatic fever may occur and must not be confused with rheumatic pneumonia Its cause is obscure but it may be connected with the long recumbent posture It is seen in many serious illnesses which confine a patient to bed for a long time e.g. typhoid fever Sometimes of course collapse of the left lower lobe may be due to pericardial effusion or to a greatly dilated heart Pulmonary congestion or œdema and infarcts of the lung should be recognised without difficulty

7 *Tolerance to Heparin* Patients with acute rheumatic fever show a remarkable tolerance to heparin and possibly to other sulphated polysaccharides This is at present under investigation and may prove a useful test for the active rheumatic state (Abrahams 1949)

EVIDENCE OF CARDITIS

To establish the diagnosis of rheumatic carditis may be one of the most difficult clinical problems in medicine There are however six reliable signs when active rheumatism can be diagnosed on other grounds

1 *Pericarditis* Transient or more persistent pericardial friction with or without effusion can be recognised in about 10 per cent of cases Rheumatic pericarditis has no special features except that it is acute often fails to alter the electrocardiogram and leaves no important sequelæ Effusion is common samples are straw coloured sterile and have the physical properties of an exudate The effect of salicylates is doubtful Paracentesis is only required to relieve cardiac compression or breathlessness due to collapse of the lung secondary to very large effusions Clinical details are similar to those in other forms of pericarditis and are described on page 341

2 *Cardiac enlargement* Displacement of the apex beat to the left is common in active rheumatic carditis but not always easy to interpret Any of the usual causes of cardiac displacement may be responsible particularly collapse or partial collapse of the left lower lobe, and abdominal distension The size of the heart is best determined by means of serial skiagrams (fig 806) even then enlargement is often a matter of opinion Again appreciable dilatation of the heart may occur in any fever natural or artificial

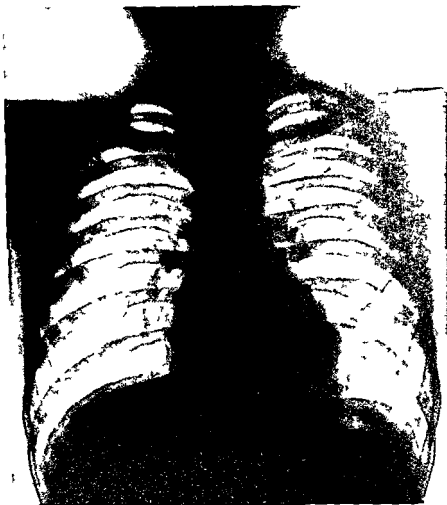


Fig 8 06 (a)—29th October 1948

Serial ski grams showing rapid development of cardiac enlargement as a result of active carditis

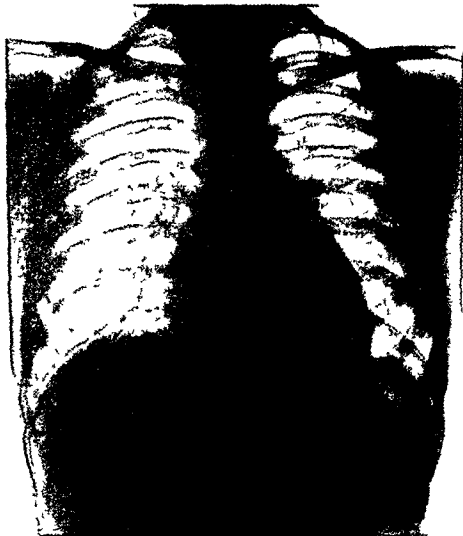


Fig 8 06 (b)—30th November 1948

Serial skiagraphs showing rapid development of cardiac enlargement as a result of active carditis



Fig 8 06 (c)—9th December 1948

Serial skiagrams showing rapid development of cardiac enlargement as a result of active
c r d i t s

(Weens and Heyman 1946) possibly because the cardiac output is raised. Unmistakable cardiac enlargement in a child with rheumatic fever is usually due to established valve lesions or to heart failure if those are absent however it must be accepted as evidence of carditis.

3 *Heart failure* The development of congestive heart failure may be accepted as proof of carditis in a patient with active rheumatism. The jugular venous pressure then rises the liver distends and the cardiac output falls. Oedema is uncommon. In otherwise uncomplicated cases orthopnoea paroxysmal cardiac dyspnoea and pulmonary oedema only occur when there is advanced aortic or mitral valve disease.

4 *Mitral diastolic murmur* The best evidence of carditis is the development of a soft mitral diastolic murmur (Carey Coombs murmur) in the absence of other signs of mitral stenosis. Autopsies have confirmed the fact that this murmur may occur in active rheumatic carditis when the mitral valve is scarcely altered (Bland White and Jones 1935). It has therefore been suggested that its mechanism depends upon left ventricular dilatation that it is related to the Austin Flint murmur and to the soft mitral diastolic murmur which is occasionally heard in hypertensive or thyrotoxic heart failure. But it must be pointed out that the Carey Coombs murmur is frequently heard when no enlargement of the heart can be demonstrated and it is more reasonable to believe that some change in the structure of the mitral valve is responsible. Whatever the explanation there is no doubt that this murmur occurs early in the course of rheumatic carditis and may disappear as activity subsides. More frequently however it is more or less persistent or reappears on the least provocation until pre systolic accentuation proclaims the development of mitral stenosis (Carey Coombs 19-4). In a series at Taplow it was transient in 20 per cent (Wood 1949).

5 *Aortic diastolic murmur* The development of an aortic diastolic murmur is equally good evidence of active carditis if its previous absence can be guaranteed. No other signs of aortic incompetence may be present be

cause the initial leak is small. Like the Carey Coombs murmur a basal diastolic murmur is occasionally transient (10 per cent). When first heard it may be remarkably short and its onset very slightly delayed perhaps not developing until the left ventricular pressure has fallen below zero at the end of isometric relaxation.

6 *Electrocardiographic abnormalities* Obvious electrocardiographic changes are the exception rather than the rule.

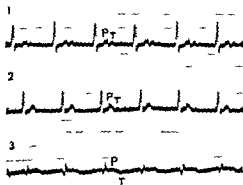


FIG. 807.—Electrocardiogram showing prolongation of the P-R interval in a case of active rheumatic carditis.

thus an apparently normal graph in no way excludes carditis. Serial records however may show transient prolongation of the P R interval in about 10 per cent of cases (fig 8 07). Normal conduction may be restored in 90 per cent of such cases by means of 1 to 3 mg. of atropine sulphate.

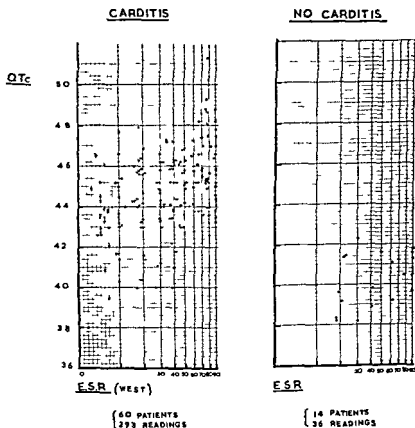


Fig 8 08—Graph showing QT plotted against the sedimentation rate in 60 cases of active rheumatic carditis and in 14 rheumatic fever controls without carditis.
(By I. F. D. Derksham)

(Bruenn 1937) Occasionally partial heart block is persistent or progresses to complete block. Nodal rhythm is sometimes seen and rarely bundle branch block. Alterations of the R S T segment usually denote pericarditis and exhibit the characteristic T pattern.

Prolongation of the Q T interval was found by Taran and Szilagyi (1947) in practically all cases of active carditis and this has been confirmed by Abrahams (1949) at the rheumatic fever centre at Taplow. In fig 8 08 the corrected Q T interval (QT_c) in sixty cases of active carditis and fourteen

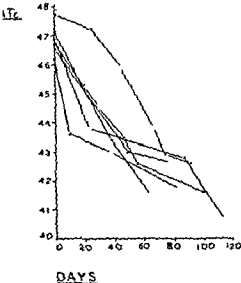


Fig 809—Behaviour of QT in six cases of acute rheumatic carditis with rapid clinical recovery

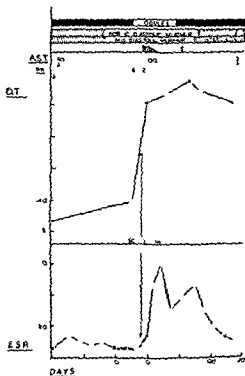


Fig 810—Prolonged QT following a recurrence of active rheumatic carditis

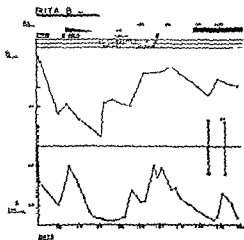


Fig 811—Graph showing a lapse due to getting up when QT was a bit grossly prolonged

(By courtesy of Dr Derek Abraham)

cases of rheumatic fever without carditis has been plotted against the sedimentation rate on a logarithmic scale. It will be seen that about 90 per cent of those with carditis have a $Q T_c$ longer than 0.42 second whereas all but one of those without carditis fall below this level. Fig. 8.09 shows the behaviour of $Q T_c$ in six typical cases of acute rheumatic carditis with rapid recovery. In fig. 8.10 a relapse is portrayed; it may be observed that $Q T_c$ then remains grossly prolonged although the sedimentation rate is falling towards normal. If the long $Q T_c$ is ignored and the patient is allowed up, relapse is likely to occur (fig. 8.11).

Paroxysmal tachycardia may occur but contributes little to establishing an etiological diagnosis; on the other hand auricular fibrillation or flutter developing in the course of active rheumatism provides good evidence of carditis although there is usually old standing mitral stenosis as well.

7 *Gallop rhythm* Gallop rhythm is more inconclusive; if it is presystolic carditis is probable; if it is proto diastolic it cannot be distinguished clinically from a normal third heart sound; if it is a summation gallop no conclusion can be drawn though it is suspicious. Analysis may be difficult for undue tachycardia is common in these cases whilst carotid sinus slowing is minimal.

8 *Systolic murmurs* Finally there is the problem of the apical systolic murmur. May it be disregarded or does it indicate mitral valvulitis? Considerable experience is required to gauge its significance. The systolic murmur of mitral incompetence associated with active rheumatic carditis may be due to ring dilatation or to mitral valvulitis. It usually begins with the first heart sound and lasts throughout systole; in quality it is commonly loud, smooth and blowing. Sometimes however the murmur may be rough or musical and may be accompanied by a thrill. When such murmurs offer the only evidence of carditis in children with rheumatic fever or chorea chronic rheumatic heart disease develops subsequently in 45 per cent of cases; under similar circumstances when the murmur is soft and unimpressive subsequent rheumatic heart disease develops in only about 9 per cent (Boone and Levine 1938; Kuttner and Markowitz 1948).

It will be seen then that the diagnosis of active rheumatic carditis rests upon six features; the presence of any one of which is diagnostic *when associated with active rheumatism*. Gallop rhythm and a mitral systolic murmur properly analysed and interpreted may be suggestive but are rarely conclusive.

When there is no extra cardiac evidence of active rheumatism the problem is different. Under these circumstances pericarditis may be tuberculous or pyogenic; cardiac enlargement and failure may be due to other causes; mitral and aortic diastolic murmurs may be attributed to old valve lesions and electrocardiographic abnormalities may have other explanations. The differential diagnosis may then include congenital heart disease, tuberculous or pyogenic pericarditis, Pick's disease, subacute bacterial endocarditis, diphtheritic and other forms of carditis, pulmonary heart

disease and anæmia. Nevertheless rheumatic carditis may occur without evidence of rheumatism elsewhere and active rheumatism may be afebrile and associated with a normal white count and E S R. A diagnosis of rheumatic carditis is therefore justified if the cardiac lesion is characteristic even when there is no other evidence of rheumatism or even of an inflammatory state. Again if there is evidence of an inflammatory state a suitable cardiac lesion may proclaim its rheumatic nature when there is no other rheumatic manifestation.

COURSE AND TREATMENT

All cases of acute or subacute rheumatic polyarthritides should be put to bed and treated with sodium salicylate 15 to 20 grains (1 to 1.5 G) combined or not with twice as much sodium bicarbonate three hourly (Lees 1904) until relieved or until ringing in the ears and deafness pronounce intoxication when the interval between doses may be increased to four hours. The effective blood level of salicylates is about 30 mg per cent. Fever, pain and joint effusions usually subside quickly but proliferative lesions including carditis are resistant. The action of salicylates is uncertain it has been suggested that they may inhibit antibody formation (Derick Hitchcock and Swift 1927-8). Toxic effects are minimised by alkalis this has been attributed to an increased rate of salicylate excretion in their presence (Parker 1947 1948). Peters (1947) however did not find that this materially affected the blood level and claimed that alkalis enhanced the therapeutic benefit of salicylates. Toxic effects include petechiæ associated with prolongation of the prothrombin time (Link *et al* 1943). Fatal hæmorrhagic encephalopathy has been reported (Ashworth and McKemie 1944). Circulating prothrombin can usually be restored by means of vitamin K in doses of 10 to 50 mg (Shapiro 1944).

When the patient has been free from symptoms for a week or if he fails to derive benefit salicylates should be stopped. If clinical relapse follows no harm is done for the exudative lesion is relatively innocent and is soon controlled by another course.

By far the best index of activity is the E S R which should be measured weekly and as a rule the patient should not be allowed up until it is normal. *This applies especially to children in whom carditis should be assumed for purposes of early management.* Adults without evidence of previous or present carditis may be treated more leniently and may be allowed up as soon as they appear well enough on clinical grounds. The duration of bed rest varies between a week or two and several months according to the severity and persistence of the active process. If patients are allowed up too soon swift relapse is the rule.

Joints may be partly or completely immobilised in the position of maximum comfort in the acute phase and some soothing lotion such as oil of wintergreen or lead and opium may be applied. Contractures are not a

feature of rheumatic fever but foot drop should be prevented and elbows and knees should not be immobilised too long

Recent work at the Mayo Clinic on the beneficial effect of compound E (17 hydroxy 11 dehydrocortico sterone) on rheumatoid arthritis (Hench *et al* 1949) has provided a new approach to the treatment of rheumatic states in general but it is too early to comment on the results so far obtained in rheumatic fever. Joint pains appear to be relieved as quickly as with salicylates but a beneficial effect on carditis has yet to be demonstrated. Cortisone (adrenal cortical hormone) is the active agent the liberation of increased quantities of natural cortisone may be brought about by the administration of A C T H (adrenocorticotrophic hormone of the pituitary)

Sulphonamides and penicillin are of no value except perhaps in the treatment of the provocative streptococcal infection or if the latter persists. Tonsillectomy is only necessary if there is chronic sepsis or if there is recurrent tonsillitis it has little influence on the disease and does not prevent relapse or recurrence (Ash 1938). A good nourishing diet fresh air vitamins especially vitamin C appropriate treatment of secondary anaemia and high morale are more important.

Chorea usually lasts 6 to 12 weeks. Patients should be put to bed during the active phase and may need heavy sedation. If there is no evidence of carditis they may be allowed up when recovery begins. They should be kept away from school and from social engagements until well.

Carditis requires absolute rest. Little else is of lasting value. Digitalis is helpful when there is congestive heart failure and although the therapeutic dose is said to be close to the toxic all effects have not been observed at Taplow. Mercurial diuretics and a low sodium diet are rarely necessary.

Absolute rest means that the patient is allowed to do nothing for himself he is washed and fed and must use bed pan and urine bottle. Diet should be light and constipation avoided. In the past it was usual to insist on nursing the patient in the horizontal position with one low pillow but it is clear from experience gained in the treatment of angina decubitus and of paroxysmal cardiac dyspnoea and from certain direct investigations in man that the cardiac output and therefore the work of the heart is greater in the horizontal than in the upright position owing to the influence of gravity on the venous filling pressure. It is therefore logical to nurse patients with carditis in the sitting posture. The wisest course may be to choose the position of maximum comfort whether lying or sitting unless there is failure when the latter should be insisted upon.

Convalescence from carditis should be extended over several months the regime being similar to that for pulmonary tuberculosis. Relapse is common and may be due to over exertion exposure emotional upset cold damp weather and to almost any infection. Relapse follows the advent of the responsible agent immediately and must be distinguished from a recurrence or second attack of rheumatic fever in which streptococcal infection

is always to blame, and following which a latent interval can usually be recognised. To avoid such infection prophylactic sulphonamide may be given in doses of 1 to 2 G daily whenever some risk is encountered. At least one recurrence occurs in two thirds of all cases usually within three years (Roth, Lingg and Whittemore 1937).

It is as important to prevent cardiac neurosis in patients with organic heart disease as it is in those without. This is a difficult task in susceptible individuals for reassurance cannot very well be unconditional. Rheumatic carditis may be symptom free and pass without influencing the subject's activities at all. Thus only about 55 per cent of cases of mitral stenosis give a history of the original attack (Parkinson and Hartley 1946). Many others are only restricted by subacute rheumatism. Little immediate harm comes to these patients indeed there is no direct evidence that subsequent development of mitral stenosis could have been prevented by bed rest at the time of active inflammation. It follows that failure to diagnose carditis when it is present in rheumatic fever is not necessarily disastrous. On the other hand its diagnosis in error may not be far short of it for the resulting cardiac neurosis which is so common may be life long and may be more incapacitating than organic heart disease. Physicians should be more aware of their responsibility in this respect. Too much emphasis is laid on overlooking a mild lesion not enough on finding what is not there. The most common mistake is to misinterpret tachycardia. A patient confined strictly to bed for several weeks with rheumatic fever is fully aware that his heart may be involved and is likely to become nervous on that account. Tachycardia may then be due to anxiety. Again the autonomic nervous system is frequently disturbed by fever and infections of all kinds tachycardia, dizziness, headache and fatigue may result especially during convalescence when activities are resumed. Such findings call for reassurance and rehabilitation not for alarm and further rest.

In the absence of diagnostic evidence of carditis throughout the active phase of rheumatic fever subsequent medical management should be based on the assumption that none existed not upon the fear that it escaped recognition and patients should be sent for convalescence as after any other fever of equal severity. This attitude is based not on the belief that carditis does not occur in a certain percentage of children with rheumatic fever but on the fact that if it does occur in an undetectable degree it is either of no consequence or it is not aggravated by this kind of management and on the fact that the over cautious attitude breeds neurosis.

MORTALITY

The mortality rate from active rheumatic carditis in childhood and adolescence is about 6.5 per cent (Scott 1943). About half the fatalities have some important complication. In the majority (62 per cent) death occurs within five years of the initial attack (Bland and Jones 1938). Bad

prognostic omens include rapid cardiac enlargement obvious nodules pericarditis auricular fibrillation and congestive heart failure Sudden unexpected death is rare in rheumatic carditis in contrast to its frequency in diphtheritic and other forms of toxic carditis There were only three such instances among a group of 7165 cases of active rheumatic fever reported by Griffith and Huntington (1946) coronary angitis was blamed

REFERENCES

- Abrahams D G (1949) 'The Q T interval in acute rheumatic carditis' *Brit Heart J* 11 342
- Aschoff L (1904) Zur Myocarditisfrage *Verhandl d deutsch path Gesellsch* 8 46
- Ash R (1938) Influence of tonsillectomy on rheumatic infection *Amer J Dis Child* 55 63
- Ashworth C T and McKernie J I (1944) Haemorrhagic complications with death probably from salicylate therapy report of 2 cases *J Amer med Ass* 126 806
- Bach F Hill N G Preston T W and Thornton C E (1939) Juvenile rheumatism in London *Ann rheum Dis* 1 210
- Barlow T and Warner F (1881) On subcutaneous nodules connected with fibrous structures occurring in children the subjects of rheumatism and chorea *Trans internat Med Cong* 4 16 London
- Bland E F and Jones T D (1938) Fatal rheumatic fever *Arch int Med* 61 161 — White P D and Jones T D (1935) 'The development of mitral stenosis in young people with a discussion of the frequent misinterpretation of a mid-diastolic murmur at the cardiac apex' *Amer Heart J* 10 905
- Boone J A and Levine S A (1938) 'The prognosis in potential rheumatic heart disease and rheumatic mitral insufficiency' *Amer J med Sc* 195 764
- Bradley W H (1932) Epidemic of acute rheumatism in public school *Quart J Med* 1 79
- Bruenn H G (1937) Mechanism of impaired auriculo ventricular conduction in acute rheumatic fever' *Amer Heart J* 13 413
- Bywaters E G L (1950) 'The relation between heart and joint disease including Rheumatoid heart disease and Chronic post rheumatic arthritis (Type Jaccoud)' *Brit Heart J* 12 101
- Cavelti P A (1947) Studies on the pathogenesis of rheumatic fever II Cardiac lesions produced in rats by means of autoantibodies *Arch Path* 44 13
- Cheadle W B (1889) Rheumatic state in childhood London
- Coburn A F (1931) Factor of infection in the rheumatic state Baltimore p 35 — (1933) Haemorrhagic manifestations of rheumatic fever *Amer J Dis Child* 45 933 — and Pauli R H (1935) Studies on immune response of rheumatic subject and its relationship to activity of rheumatic process active and passive immunization to hemolytic streptococcus in relation to rheumatic process *J clin Invest* 14 755
- Collis W R F (1931) Acute rheumatism and hemolytic streptococci *Lancet* 1 1341
- Coombs C F (1924) Rheumatic heart disease Bristol p 203
- Dawson M H Olmstead M and Boots R H (1932) Agglutination reactions in rheumatoid arthritis Agglutination reaction with streptococcus hemolyticus *J Immunol* 23 187 205
- Derick C L Hitchcock C H and Swift H F (1947-8) The effect of anti-rheumatic drugs on the arthritis and immune body production in serum disease *J clin Invest* 5 427
- Eiman J and Gouley B A (1948) Rheumatic pneumonitis *Arch Path* 5 558

is always to blame, and so losing which a latent interval can usually be recognised. To avoid such infection, prophylactic sulphonamide may be given in doses of 1 to 2 G daily whenever some risk is encountered. At least one recurrence occurs in two-thirds of all cases usually within three years (Roth, Lingg and Whittemore 1937).

It is as important to prevent cardiac neurosis in patients with organic heart disease as it is in those without. This is a difficult task in susceptible individuals, for reassurance cannot very well be unconditional. Rheumatic carditis may be symptom free, and pass without influencing the subject's activities at all. Thus only about 5 per cent of cases of mitral stenosis give a history of the original attack (Parkinson and Hardley 1946). Many others are only restricted by subacute rheumatism. Little immediate harm comes to these patients; indeed there is no direct evidence that subsequent development of mitral stenosis could have been prevented by bed rest at the time of active inflammation. It follows that failure to diagnose carditis when it is present in rheumatic fever is not necessarily disastrous. On the other hand, its diagnosis in error may not be far short of it, for the resulting cardiac neurosis, which is so common, may be life-long and may be more incapacitating than organic heart disease. Physicians should be more aware of their responsibility in this respect. Too much emphasis is laid on overlooking a mild lesion, not enough on finding what is not there. The most common mistake is to mistakingly expect tachycardia. A patient confined strictly to bed for several weeks with rheumatic fever is fully aware that his heart may be involved and is likely to become nervous on this account. Tachycardia may then be due to anxiety. Again, the autonomic nervous system is frequently disturbed by fever and infections of all kinds. Tachycardia, dizziness, headache and fatigue may result especially during convalescence when activities are resumed. Such findings call for reassurance and rehabilitation, not for alarm and further rest.

In the absence of diagnostic evidence of carditis throughout the acute phase of rheumatic fever subsequent medical management should be based on the assumption that none existed, not upon the fear that it escaped recognition and patients should be sent for convalescence as after any other fever of equal severity. This attitude is based, not on the belief that carditis does not occur in a certain percentage of children with rheumatic fever but on the fact that if it does occur to an undetectable degree, it is either of no consequence or it is not aggravated by this kind of management, and on the fact that the over-cautious attitude breeds neurosis.

MORTALITY

The mortality rate from active rheumatic carditis in childhood and adolescence is about 6.5 per cent (Scott, 1943). About half the fatalities have some important complication. In the majority (62 per cent) death occurs within five years of the initial attack (Bland and Jones 1936). End

- Paul J R (1928) Pleural and Pulmonary lesions in rheumatic fever *Medicine* 7 383
- Perry C B (1937) Erythema marginatum (rheumatism) *Arch Dis Child* 12 233
- Peters J T (1947) The necessity and possibility of giving detoxified large oral doses of salicylates in the treatment of rheumatic fever in order to prevent or cure the inflammatory stage of carditis *Acta med Scand* 128 51
- Poynton F J and Paine A (1900) The etiology of rheumatic fever *Lancet* ii 861 — (1913) *Researches on rheumatism* London
- Rich A R and Gregory J E (1943) On the anaphylactic nature of rheumatic pneumonitis *Bull Johns Hopk Hosp* 73 465 — (1943) Experimental evidence that lesions with the basic characteristics of rheumatic carditis can result from anaphylactic hypersensitivity *Ibid* 73 239 — (1944) Further experimental cardiac lesions of the rheumatic type produced by anaphylactic hypersensitivity *Ibid* 75 115
- Ritchie W T (1939) Acute rheumatic carditis *Lancet* ii 582
- Ritter S A, Gross L and Kugel M A (1927-8) Blood vessels in the valves of normal human hearts *Amer Heart J* 3 433
- Rogen A S (1947) The heart in rheumatoid arthritis *Brit med J* i 87
- Roth I R, Lingg C and Whittemore A (1937) Heart disease in children rheumatic group certain aspects of age at onset and of recurrences in 488 cases of juvenile rheumatism ushered in by major clinical manifestations *Amer Heart J* 13 36
- Schlesinger B (1930) The relationship of throat infection to acute rheumatism in childhood *Arch Dis Child* 5 411
- Scott G E M (1943) Rheumatic infection in childhood survey from Children's Hospital Melbourne with addendum on follow up system of almoners of the hospital *Med J Australia* 2 309
- Shapiro S (1944) Studies on prothrombin effect of synthetic vitamin K on prothrombinopenia induced by salicylate in man *J Amer med Ass* 125 546
- Shaw A F B (1929) Topography and pathogenesis of lesions in rheumatic fever *Arch Dis Child* 4 155
- Sheldon W (1931) On acute rheumatism following tonsillitis *Lancet* i 1337
- Stuart Harris C H (1935) A study of hæmolytic streptococcal fibrinolysis in chronic arthritis rheumatic fever and scarlet fever *Ibid* ii 1456
- Sutton L P and Dodge K G (1938) The relationship of Sydenham's chorea to other rheumatic manifestations *Amer J med Sci* 195 656
- Taran L M and Szilagyi N (1947) The duration of the electrical systole (QT) in acute rheumatic carditis in children *Amer Heart J* 33 14
- Todd E W (193-) Anti-hæmolytic titres in hæmolytic streptococcal infections and their significance in rheumatic fever *Brit J exper Path* 13 248
- Wearn J T, Bromer A W and Zachiesche L J (1936) The incidence of blood vessels in human heart valves *Amer Heart J* 2 —
- Weens H S and Heyman A (1946) Cardiac enlargement in fever therapy induced by intravenous injection of typhoid vaccine *Arch int Med* 77 307
- Werner M (1938) Über die Ursachen der Verquellung der kollagenen Fasern bei hyperergischen Entzündung (Arthussches Phänomen) (Zugleich ein Beitrag zur Funktion des Bindegewebes) *Virchows Arch f path Anat* 301 55
- Wilson M G (1940) *Rheumatic fever* New York
- Wood P H (1949) Cardiac complications of rheumatic fever *Proc Roy Soc Med* 43 195

CHAPTER IX

INACTIVE RHEUMATIC HEART DISEASE

RHEUMATIC carditis refers to active inflammation of the heart. The after-effects, which include valvular sclerosis, patchy myocardial fibrosis and adherent pericardium are best described under the general heading of rheumatic heart disease to which the appropriate anatomical abnormality may be appended. Thus we may speak of rheumatic heart disease with mitral stenosis.

Incidence About 5 per cent of healthy young adults give a previous history of rheumatic fever in childhood (Parkinson and Hartley 1946). It is clear therefore that all patients who have rheumatic fever do not later develop clinical rheumatic heart disease. According to Carey Coombs (1924) 50 per cent of children who have their first attack of rheumatic fever before they are five years old and 25 per cent of those whose first attack occurs after the age of ten subsequently develop rheumatic heart disease.

From large samples of the younger male population of Great Britain examined for military service between 1939 and 1945 it was calculated that there were about 240 000 cases of rheumatic heart disease of both sexes between the ages of 18 and 44 in Great Britain at that time or about 7 per cent of the population in that age-group (Parkinson 1945). Rheumatic heart disease accounts for approximately 25 per cent of all cases of heart disease in temperate climates and causes about 16 000 deaths annually in England and Wales.

Practically all clinical cases of inactive rheumatic heart disease have one or more valve lesions. The mitral valve is involved in 85 per cent, the aortic in 44 per cent, the tricuspid in 10 to 16 per cent and the pulmonary in 1 to 2 per cent (Cabot 1926). Mitral disease is more common in females in the ratio of 3 : 2, aortic valve disease is equally more common in males.

LESIONS OF THE VALVES

1. MITRAL INCOMPETENCE

In the last century this was the most common valve lesion diagnosed. Owing to the exertions of Mackenzie, Lewis, Parkinson and others recent years have witnessed a diagnostic revolution so that now a physician who asserts that a patient has organic mitral incompetence must be very sure of his grounds. The change in outlook has saved a host of normal subjects from invalidism. Nevertheless mitral incompetence is a real entity and the

whole problem must be critically examined. The difficulty lies in the interpretation of apical systolic murmurs. In the past these were nearly always attributed to mitral incompetence and the latter was believed to be the consequence of valvular disease. It is now known however that apical systolic murmurs may be cardiac or extra cardiac and if cardiac may be due to mitral incompetence or to some other factor. Mitral incompetence itself may be organic or functional. These possibilities will be considered in detail.

EXTRA CARDIAC MURMURS

The heart expanding in diastole may press upon a portion of lung so that the latter suddenly decompressed in cardiac systole sucks in air and gives

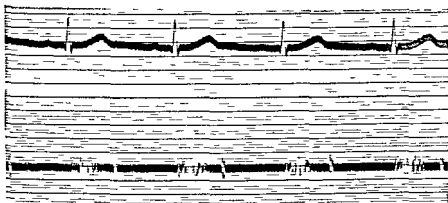


Fig 9 01—Phonocardiogram showing an innocent systolic murmur at the mitral area

(By out y f D l b e y L tham)

rise to a systolic vesicular murmur. This is soft and varies with respiration and with change of posture. Normal friction between two healthy serous membrane adequately lubricated may give rise to a murmur indistinguishable in quality from the ordinary soft apical systolic bruit (Ortiz 1933). Such murmurs are probably exaggerated by overaction of the heart as in thyrotoxicosis, anemia and the anxiety states.

Innocent systolic murmurs have been said to begin some time after the first heart sound (Evans 1947) but this is not generally accepted. Many such murmurs beginning early in systole (fig 9 01).

CARDIAC MURMURS

(a) *Without mitral incompetence* In aortic stenosis a systolic murmur undoubtedly produced at the aortic valve is sometimes heard best at the apex beat. It follows that a functional murmur arising at the base may also be heard best at the apex beat. It should be understood that this site commonly represents the apex of the left ventricle and is therefore in direct

communication with the aorta. Functional murmurs may result from turbulence at the root of the aorta in normal subjects especially when the cardiac output is increased. Harsh or musical apical systolic murmurs may occur in hearts which offer no obvious mechanical explanation when examined later after death. It is thought that these may be due to unusual vibration of the chordæ tendineæ.

(b) *Mitral incompetence (functional)* Experiments after death show that the aortic valve is competent and the mitral incompetent when water is poured into the aorta or left ventricle respectively. During life proper mitral closure depends upon adequate constriction of the mitral ring and upon sufficient slackness of the inelastic chordæ tendineæ which, acting as guy ropes anchor the valve cusps to the wall of the left ventricle. When that chamber is dilated the mitral ring is expanded and stretched chordæ tendineæ may prevent the cusps falling back sufficiently to come into close apposition. Functional mitral incompetence in left ventricular failure is therefore more or less inevitable. It may also complicate left ventricular dilatation in the hyperkinetic circulatory states and in any form of carditis. Functional mitral incompetence of this kind seems to occur naturally in some otherwise normal hearts perhaps due to some slight fault in architecture. It is of no consequence.

(c) *Mitral incompetence (organic)* Organic mitral incompetence refers to disease of the mitral valve which prevents its proper closure. It is nearly always rheumatic and may be associated with shortening and thickening of the chordæ tendineæ. In the great majority of cases the mitral ring and base of the cusps are also sclerosed resulting in mitral stenosis. In clinical diagnosis the latter is so much the more important that the other is hardly worth mentioning in its company. But if there is no stenosis organic mitral incompetence leads to a well defined clinical picture. During systole the left auricular pressure and volume rise well beyond normal limits and the chamber expands visibly. During diastole the left ventricle becomes abnormally distended as a result of this high filling pressure. The stroke volume propelled into the aorta thus remains normal. In other words increased distension of the left ventricle compensates for the leak. In time both the left auricle and ventricle may become considerably enlarged.

The diagnosis of organic mitral incompetence is based upon the following features

1. An impressive mitral systolic murmur. This is not necessarily loud or harsh but it usually begins early, lasts throughout systole and has a characteristic blowing quality (fig. 9.02a). The intensity of the murmur should not be altogether disregarded as follow up studies on children with suspected rheumatic carditis have demonstrated clearly that those with loud murmurs are five times more likely to develop chronic rheumatic heart disease than those with soft murmurs (Boone and Levine 1938, Kuttner and Markowitz 1948). The loud late systolic murmur which seems to increase in intensity as it merges into the second heart sound

(fig 9 02b) though sometimes innocent may undoubtedly be due to organic mitral incompetence thus it may be associated with mitral stenosis or with expansile pulsation of the left auricle

2 Mitral systolic thrill As a general rule a thrill means organic disease but there are exceptions. Conversely many cases of mitral incompetence

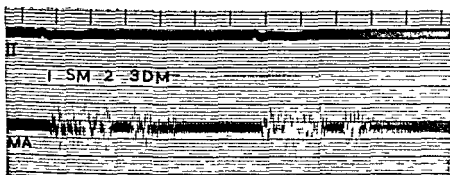


Fig 902 (a)—Phonocardiogram showing a systolic murmur due to organic mitral incompetence

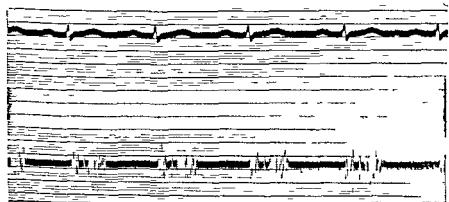


Fig. 902 (b)—Loud late systolic murmur in a case of organic mitral incompetence.

(R) *ow t y f D H H m E d A bey L (hau)*

have no thrill. When there is aneurysmal dilatation of the left auricle the thrill may be felt to the right of the sternum.

3 Left ventricular enlargement This is variable. It must be judged on clinical, radiological and electrocardiographic evidence (fig 9.03).

4 Dilatation and systolic expansion of the left auricle. Skiagrams showing left auricular dilatation in the antero posterior or right anterior oblique position do not distinguish mitral incompetence from mitral stenosis (fig.

233) but when viewed fluoroscopically the former is recognised by expansile pulsation of the left auricle during ventricular systole. If cardiac movements are correctly interpreted this is diagnostic. Kymography facilitates analysis.

There are two distinct clinical types of organic mitral incompetence: those that develop aneurysmal dilatation of the left auricle and that behave like cases of mitral stenosis; and those that develop increasing left ventricular enlargement and which

behave more like cases of aortic stenosis (with which they are readily confused).

The prognosis of organic mitral incompetence is fair: cases with aneurysmal dilatation of the left auricle usually die between the ages of 30 and 45; those with large left ventricles may be expected to survive to the age of 50 or so but then succumb rather suddenly to irreversible heart failure.



Fig. 903.—Skiagram showing dilatation of the left ventricle in a case of organic mitral incompetence. On fluoroscopy the outline of the left auricle expanded visibly during systole.

2 MITRAL STENOSIS

Pathology. Reparative fibroblastic proliferation already described as part of the rheumatic lesion has unfortunate results when situated at the base of the mitral cusps or in the mitral ring; for here subsequent contracture leads to

stenosis of the valve. This is the common sequel of rheumatic carditis as the base of the valve is its most vascular part and therefore the most effected. The lesion takes at least two years to develop sufficiently to give rise to physical signs, usually much longer (Carey Coombs, 19-4).

The degree of narrowing varies from an amount insufficient to have any effect on function to tight button-hole stenosis with an orifice which barely admits a pencil. Calcification is common in elderly subjects. Associated mitral incompetence is the rule but is relatively unimportant.

Disturbance of function. The blood flow being obstructed at the mitral valve the pressure rises in the left auricle and pulmonary veins. This at first compensates for the obstruction and insures full left ventricular filling; the greater speed of blood flow through the mitral orifice making up for its smaller cross section.

Next the increased pressure in the left auricle and pulmonary veins

causes a rise in pulmonary arterial pressure so that the normal pressure gradient is maintained. There is little to support the view that pulmonary hypertension in mitral stenosis does not depend on the left auricular pressure but upon some hypothetical reflex or upon rheumatic pulmonary arteritis (Gouley 1938). On the contrary left auricular pressures when measured at thoracotomy may be very high indeed and amply explain mean pulmonary artery pressures over 30 mm Hg.

When relatively healthy the left auricle hypertrophies and so increases the presystolic pressure gradient between auricle and ventricle. This contributes further to adequate left ventricular filling. Not infrequently however myocardial fibrosis weakens the left auricle and favours dilatation of that chamber. Sooner or later left ventricular filling is hampered and the cardiac output tends to fall or cannot meet the demands of effort. The pulse volume becomes small and the aorta and left ventricle hypoplastic.

At the same time the right ventricle hypertrophies to cope with pulmonary hypertension and the pulmonary artery dilates. There is a permanent shift in blood distribution more being held in the pulmonary circulation and left auricle than usual and correspondingly less on the systemic side. The vital capacity and lung volume are thus reduced and the pulmonary circulation time prolonged. If the blood volume does not increase compensatory systemic vasoconstriction results adapting the systemic vascular capacity to its reduced contents and so maintains the blood pressure. Hence the small pulse is firm and the skin is cold, pale and cyanotic.

Dyspnoea at this stage is not due to an increase of carbon dioxide acting on the respiratory centre nor to reduced oxygen tension acting on chemoreceptors in the carotid sinus for the gaseous exchange is still normal. It is intimately connected with the engorged pulmonary circulation. This excites a respiratory reflex through pulmonary vagal afferents whereby breathing becomes quicker (Harrison 1935). The Hering Breuer reflex is ordinarily concerned with limiting the depth of inspiration and is initiated by stretch receptors which are probably situated in the walls of the alveolar ducts. The mechanism is inhibitory and is abolished by section of the pulmonary vagal afferents. Just how vascular engorgement increases the activity of the stretch receptors if these two reflexes are the same is not clear.

Likewise cyanosis is not due to reduction of the arterial oxygen saturation for this does not occur unless there is pulmonary oedema or other late changes in the lung parenchyma. Cyanosis is usually attributable to peripheral vasoconstriction.

In an attempt to increase the cardiac output the venous filling pressure tends to rise especially on effort and for a time this mechanism may prove successful but sooner or later paroxysmal cardiac dyspnoea or congestive heart failure develops.

Clinical features Well established mitral stenosis is characterised clinically

I 2

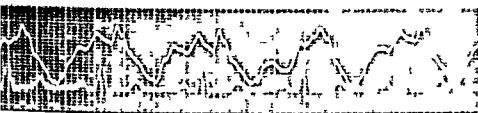


Fig 905—Phonocardiogram at the apex beat in a case of patent ductus showing a functional mitral diastolic murmur
(By courtesy of Dr Frances Gardner and Max Zobel)

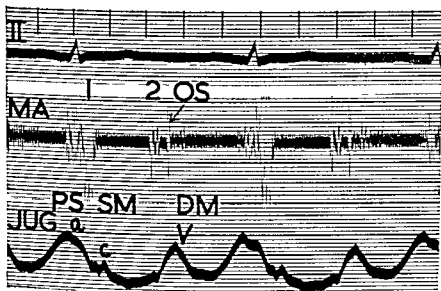


Fig 906—Phonocardiogram showing a crescendo-decrescendo murmur timed against the electrocardiogram and phlebogram
(By courtesy of Dr William Evans and Aubrey Leatham)

that continues during the period of isometric relaxation. The characteristic mitral diastolic bruit is low pitched and rumbling, is heard best with a bell stethoscope and may be exaggerated when the patient lies on the left side or when the cardiac output is increased by means of exercise or amyl nitrite. The recognition of this murmur often depends upon the examiner's ability to concentrate upon that phase in diastole in which he knows it should arise to the exclusion of all other sounds. Never was attention directed to a preconceived sound more amply rewarded. Pre-systolic accentuation (Fauvel 1843) (fig. 906) depends upon auricular contraction (Gardner 1861) and necessarily disappears when the auricles fibrillate. Occasionally functional pulmonary incompetence causes a basal diastolic murmur down the left border of the sternum—the Graham Steell murmur (Steell 1888). There is no sure way of distinguishing this murmur from that of aortic incompetence but it tends to be more local, is placed rather higher on the left and may be lower in pitch. Interpretation however is more safely based upon the presence of gross dilatation of the pulmonary artery on the one hand and upon other evidence of aortic valve disease on the other.

The mitral diastolic murmur is not quite pathognomonic of mitral stenosis for turbulence giving rise to an identical bruit may occur in active rheumatic carditis, gross aortic incompetence (Austin Flint 1862) and when left ventricular dilatation is associated with an increased pulmonary blood flow as in patent ductus arteriosus (fig. 905), ventricular septal defect, thyrotoxicosis and anaemia.

Ray appearances. Fluoroscopic reveals characteristic changes in the size and shape of the heart. The aorta is small, the pulmonary arc dilated, and the hilar vessels engorged. The dilated left auricle often causes a bump on the left border of the heart between the pulmonary artery and left ventricle, a prominence which is seen in no other condition (fig. 907). Left auricular enlargement is seen best in the right anterior oblique position with barium in the oesophagus (fig. 908).



Fig. 907.—Skiagram of a case of mitral stenosis showing dilatation of the left auricle between the pulmonary arc and the left ventricle. The left auricle may also be seen at the right border of the heart above and overlapping the right auricle.



Fig 9 08—Skigram of a case of mitral stenosis showing dilatation of the left auricle in the right anterior oblique position. The oesophagus is outlined with barium.



Fig 9 09—Skigram of a case of mitral stenosis showing dilatation of the left auricle in the left anterior oblique position.



Fig 9 10—Skigram of a case of mitral stenosis showing milary nodules in the lungs due to hæmopteroses.

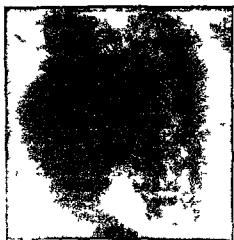


Fig 9 11—Angiocardiogram in a case of mitral stenosis showing the left auricle in the second oblique position.

As a rule the chamber enlarges backwards and to the right and may often appear in the antero posterior view just above and overlapping the right auricle (fig 9 07) Occasionally it enlarges backwards and to the left when it may be seen best in the left anterior oblique position just above and overlapping the shadow of the left ventricle (fig 9 09 and 9 11) Appearances in the lung fields occasionally resemble those of military tuberculosis (fig 9 10) They have been ascribed to pulmonary hemosiderosis similar to that seen in certain hæmolytic anæmias of childhood (Gumpert 1947)

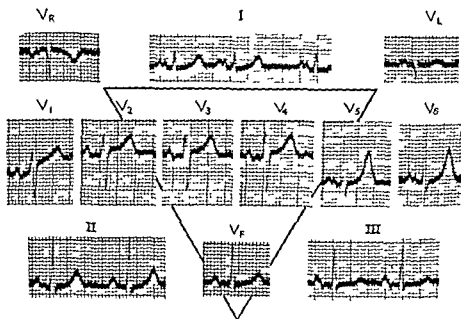


Fig 9 12—Electrocardiogram in a case of mitral stenosis showing widened bifid P waves particularly in leads I V₅ and V₆ The heart is vertical

Angiocardiograms prove that the bump on the left border of the heart between the pulmonary arc and left ventricle is the left auricle not the conus of the right ventricle

Electrocardiography Electrocardiograms commonly show widened bifid P waves particularly in leads I 2 V₄ V₅ and V₆ (fig 9 12) indicating left auricular enlargement they are not necessarily very conspicuous (fig 9 13) Occasionally P is tall and harp (fig 9 14) as in pulmonary heart disease indicating right auricular enlargement Pulmonary hypertension or tricuspid stenosis may then be responsible When there is considerable right ventricular enlargement, partial right bundle branch block is common although standard limb leads may give little indication of it (fig 9 15) Right axis deviation in mitral stenosis is usually due to a vertical heart the QR complex from the left ventricle being transmitted to lead V_F (fig 9 12)

MITRAL STENOSIS

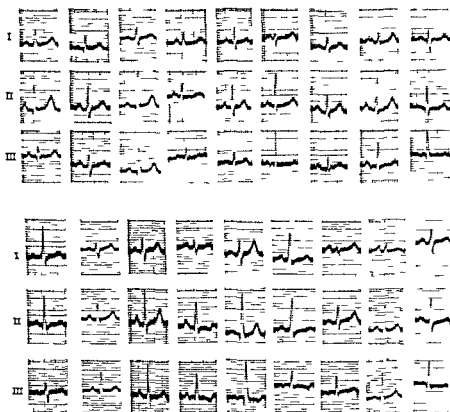


Fig 913—Electrocardiograms showing the P waves in 18 unselcted cases of mitral stenosis

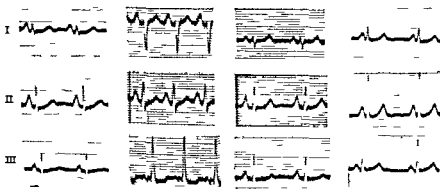


Fig 914—Standard lead electrocardiograms in 4 cases of mitral stenosis showing tall sharp P waves like those seen in pulmonary heart disease

or it may be due to partial right bundle branch block, the RS complex from V₆ being transmitted to V_L and the RSR pattern from V₁ being transmitted to V_I the heart then being electrically horizontal. In rare instances a QR complex may be seen in lead V₁ this is probably due to clockwise rotation about the longitudinal axis (viewed from below) and represents potentials at the back of the heart.

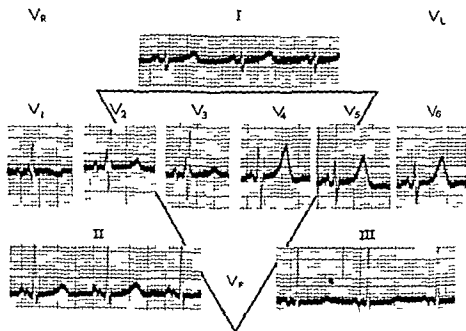


Fig 9 15—Electrocardiogram in a case of mitral stenosis showing partial right bundle branch block.

Functional studies The vital capacity and lung volume are reduced and the arm to tongue circulation time is prolonged in proportion to the degree of pulmonary congestion. The jugular venous pressure and right auricular pressure tend to be slightly elevated when there are symptoms of effort intolerance—before the development of congestive heart failure proper. When clinically the venous pressure is normal it may yet be possible to demonstrate its conspicuous elevation on effort. This is good evidence of limited cardiac reserve.

The right ventricular pressure measured by means of cardiac catheterisation is usually somewhat raised, mean figures mostly ranging between 10 and 30 mm Hg above the mean right auricular pressure. In tight stenosis with paroxysmal cardiac dyspnoea however extremely high pressures are found (30–60 mm Hg).

The arterial oxygen saturation remains normal until the terminal phase apart from attacks of pulmonary oedema. The cardiac output may be within normal limits at rest but unable to meet the demands of effort when con-

gestive failure develops it is always low. The arterio venous oxygen difference is then increased.

Complications

1 *Auricular fibrillation* Auricular fibrillation is common occurring sooner or later in 50 per cent of cases. Its incidence bears a linear relation to the age of the patient. When it occurs in adolescents or in young adults it nearly always signifies rheumatic activity. Although it may occur in paroxysms at first it soon becomes permanent and attempts to restore normal rhythm with quinidine are apt to be fruitless and dangerous. Auricular flutter is less common but by no means rare. In fact mitral stenosis is the most common cause of flutter. Such changes of rhythm are important because they may provoke heart failure and because they increase the risk of embolism.

2 *Congestive heart failure* Paroxysmal cardiac dyspnoea and pulmonary oedema are due to mitral stenosis in about 9 per cent of cases when they represent considerable obstruction at the mitral valve and a strong right ventricle acting with normal rhythm. Attacks are prone to occur on effort when the right ventricle pumps more blood into the lungs than can escape through the tight mitral orifice. They are readily induced by a rigor. Congestive heart failure may occur with normal rhythm or may follow the onset of auricular fibrillation or flutter. Recurrent rheumatic activity, intercurrent infection, pregnancy and heavy manual work are the most important provocative factors.

3 *Hæmoptysis* was first correlated with mitral stenosis by Wilson in 1830 (Rolleston 1941). There are two common causes: rupture of an engorged vessel and pulmonary infarction. Hæmorrhage from the former is apt to be short and brisk from the latter prolonged. Pulmonary apoplexy usually occurs with normal rhythm, pulmonary infarction with auricular fibrillation and heart failure.

4 *Systemic emboli* Clots form in the dilated left auricle in 12 per cent of all cases of mitral stenosis and in 24 per cent of those with auricular fibrillation (Davis and Weiss 1931). Ill advised quinidine therapy may excite the liberation of such a clot.

5 *Laryngeal palsy* Gross left auricular dilatation may compress the left recurrent laryngeal nerve and paralyse the left vocal chord (Ortner 1897). Dilatation of the left pulmonary artery assists in the process (Fetterolf and Norris 1911). Huskiness of the voice follows. The condition is not rare in advanced mitral stenosis.

6 *Collapse of the lung* Gross enlargement of the left auricle may compress the left bronchus and cause collapse of the lung (King 1838). Splaying of both bronchi may often be demonstrated radiologically.

7 *Recurrent bronchitis* Mitral stenosis appears to encourage the incidence of recurrent bronchitis. Rhonci may obscure the characteristic murmur and the correct diagnosis may then be overlooked.

8 *Bacterial endocarditis* Acute or subacute bacterial endocarditis may complicate mitral valve disease in any stage of its history including that of active rheumatism and should always be borne in mind

Associated diseases Rheumatoid arthritis (page 256) and arachnodactyly (page 255) have already been discussed. Essential hypertension was found in 28 per cent of 150 cases of mitral disease reported by Berconsky and Neuman (1945) but was regarded as fortuitous. Coincident thyrotoxicosis is not rare. Despite the fact that rheumatic fever occasionally causes coronary arteritis, angina pectoris is very uncommon in mitral stenosis and when present may be attributed to independent coronary atherosclerosis. The incidence of tuberculosis in cases of mitral stenosis is lower than in the general population.

Course and prognosis The average life history of rheumatic heart disease with mitral stenosis may be summarised as follows. The initial rheumatic attack occurs between the ages of 8 and 12. Severe cases die within ten years; these are nearly all still active. Those who make a good immediate recovery usually remain free from symptoms for about twenty years. Breathlessness on exertion sufficient to limit the patient's ordinary activities then develops and progresses to congestive heart failure within two or three years. The average age at death is about 35. Patients who die with normal rhythm tend to be younger; the average age of death for this group being 29; the average age of death in patients with auricular fibrillation is 38 (De Graff and Lingg, 1935). The duration of auricular fibrillation averages but two to three years. Such figures provide a useful basis upon which to assess prognosis in any particular case, but individual variation is great: some patients dying in adolescence, others reaching old age. Each should be judged on its own merits, due consideration being paid to effort tolerance, heart size, rhythm, congestive heart failure, and particularly to recurrent or persistent rheumatic activity. Life expectancy is little influenced by the presence of other valve lesions.

Persistent active rheumatic carditis is probably the most important factor determining prognosis. Thus De La Chapelle, Graef and Rottino (1934) demonstrated Aschoff nodes in 86 per cent of cases dying under 40 years of age, and in 33 per cent of cases over 40. Werner (1936) found activity in 66 per cent of all cases of rheumatic heart disease which had died from congestive failure.

3 AORTIC INCOMPETENCE

Pathology Rheumatic inflammation of the aortic valve may cause immediate aortic incompetence. Healing usually results in thickening, retraction and distortion of the cusps, with permanent regurgitation. In addition the cusps often become adherent to one another at their bases (fusion of the commissures) so that some degree of aortic stenosis is usual. Secondary calcification is common, especially when there is stenosis.

Effect on function The stroke volume of the left ventricle is increased by

an amount which is at least equal to the quantity of blood which leaks back during diastole. The fibres of the left ventricle become considerably stretched in diastole; the force of the heart beat is therefore augmented according to Starling's law. The initial tension is increased, isometric contraction is abbreviated, maximum pressure is higher than normal and is attained earlier in systole; the ejection phase is shortened and the pressure then falls away steeply in late systole. In other words, the shape of the pressure curve is altered so that early systole is loaded and late systole unloaded (Wiggers, 1935). The large quantity of blood pumped so quickly and powerfully into the relaxed arteries during early systole causes an abrupt percussion wave followed by late systolic collapse. The low diastolic

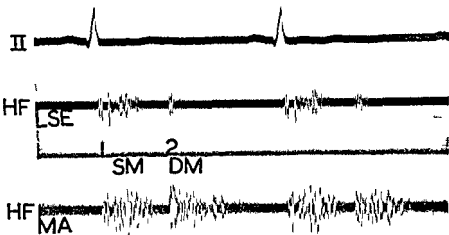


Fig. 9.16—Phonocardiogram illustrating a diminuendo aortic diastolic murmur

pressure is due partly to the aortic reflux and partly to peripheral vasodilatation; the latter encourages forward flow. Both add to the collapsing quality of the pulse.

The cardiac output per minute remains about normal or may be even a little raised, as it is in patent ductus arteriosus and arterio-venous aneurysm, which have much in common with aortic incompetence. Effort tolerance is usually remarkably good until the disease is well advanced. Sooner or later, however, left ventricular failure develops, often suddenly and unexpectedly. The heart then becomes overloaded and the output falls below normal.

Clinical features. Unlike mitral stenosis, aortic incompetence develops during the stage of active valvulitis and may be at once permanent. Its early diagnosis depends entirely upon recognising an aortic diastolic murmur heard best down the left border of the sternum and closely resembling the sound of a whispered R (Hope, 1839). In contrast to the mitral

diastolic murmur there is no gap between it and the second heart sound the one passing imperceptibly into the other. Thus the usual two beat metre of the heart sounds is not altered (fig 9 16). In distinguishing aortic from mitral diastolic murmurs the greatest stress is laid on this difference in rhythm for aortic murmurs may be heard best at the apex beat and mitral murmurs towards the base. It has already been explained that owing to the appreciable period which must elapse between the closure of the aortic and the opening of the mitral valves mitral diastolic murmurs give rise to a three beat dactylic cardiac metre.

When incompetence is more pronounced numerous changes in the heart

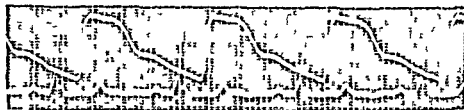


Fig 9 17—Arteriogram illustrating the water hammer pulse of aortic incompetence. The percussion wave is unusually abrupt collapse precedes the pre diastolic notch and is therefore a late systolic event.

and circulation may be recognised. Owing to enlargement of the left ventricle the apex beat is displaced downwards and to the left and the cardiac impulse is heaving. At the mitral area a diastolic murmur may develop which has all the qualities of mitral origin: there is a gap between its commencement and the second heart sound; it is soft, low pitched and rumbling; it may be accentuated in presystole. This is the Austin Flint murmur and may depend upon interference with mitral valve function by regurgitating blood or upon left ventricular dilatation. It is indistinguishable from the diastolic murmur of mitral stenosis but is rarely accompanied by a thrill.

During systole the increased volume of blood flung into the circulation raises the systolic pressure and distends the aorta and large arteries. The upstroke of the pulse wave is abrupt and of high amplitude (fig 9 17). When an artery is palpated this sudden shock feels like a water hammer (a Victorian toy consisting of a small quantity of fluid in a glass vacuum tube—Watson 1843) and on auscultation the sound heard may resemble a pistol shot.

The abrupt distension and quick collapse of large arteries is well seen in the carotids especially when the patient sits up. This characteristic visible

behaviour of an artery above heart level is Corrigan's sign (Corrigan 1832)

On auscultating the femoral or other large artery a systolic murmur is heard when the vessel is compressed when a critical pressure is applied to the artery just distal to the stethoscope a diastolic murmur may also develop. The latter was first described by Durozier (1861) whose name is attached to the sign and who attributed it to retrograde blood flow during diastole. *Durozier's sign may occur however in any condition causing a*



Fig 18—Skag am showing prominence of the aortic arch and enlargement of the left ventricle in a case of aortic incompetence



Fig 19—Second oblique position showing enlargement of the left ventricle and unfolding of the aorta

large primary pulse wave a steep predicrotic notch and a conspicuous dicrotic wave. Such an obstacle halts the blood flow at the pre dicrotic notch but is overcome by the dicrotic wave. Above the obstacle the dicrotic wave is exaggerated below it the dicrotic wave is flattened out. Hence the diastolic murmur is heard above but not below the constriction. The centrifugal direction of the passage of the wave which causes the murmur has been proved by means of simultaneous multiple phonoarteriograms (Luisada 1943).

Vasodilatation exaggerates the collapsing quality of the pulse further lowers the diastolic blood pressure and causes capillary pulsation. The latter may be demonstrated by lightly compressing a finger nail by transilluminating the tip of the finger or by pressing a glass slide against the lips. Its presence depends upon direct transmission of the arterial pulse wave to the capillaries and it occurs in any condition in which there is sufficient relaxation of the arterioles to allow this. Thus capillary pulsation

may be seen in normal subjects after a hot bath, in thyrotoxicosis, arterio-venous aneurysm fever and in most hyperkinetic circulatory states. Pulsation of the retinal veins is another common finding.

Skiagrams show enlargement of the left ventricle and prominence of the aorta. The ascending aorta pushes the superior vena cava further to the right; the aortic knob is accentuated and the descending limb appears further to the left (fig. 9 18). Unfolding of the arch is seen better in the left

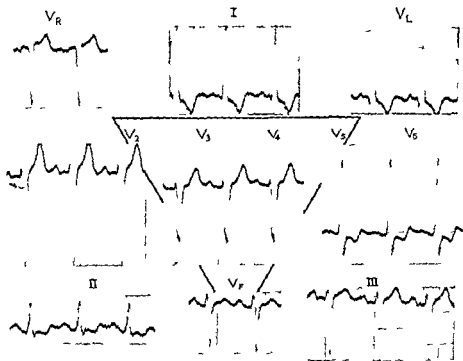


FIG. 9 20—Electrocardiogram in a case of aortic incompetence showing evidence of left ventricular enlargement

anterior oblique position (fig. 9 19). Fluoroscopy reveals exaggerated pulsation of the left ventricle and aorta. Electrocardiography may provide additional evidence of left ventricular enlargement (fig. 9 20).

Most of the features described above are common to all forms of aortic incompetence, but their degree varies according to the etiology of the lesion. Aortic incompetence may be due to a congenital bicuspid valve, to rheumatic valvulitis (active or healed), to bacterial endocarditis, syphilitic aortitis, atherosclerosis, hypertension or trauma. The special characteristics of each type are described in chapters devoted to the diseases mentioned.

In differential diagnosis, rheumatic aortic incompetence is favoured by a rheumatic history, signs of associated aortic stenosis with or without calcification, the presence of other valve lesions, absence of angina pectoris and by a normal erythrocyte sedimentation rate. It is not always easy to

be certain as to whether the mitral valve is stenosed when the chief lesion is obviously aortic incompetence for then a mitral presystolic or diastolic murmur backward displacement of the œsophagus and widened hind P waves may not have their usual significance. The practical point emerges that a case presenting as one of aortic incompetence with doubtful signs of mitral stenosis is better judged rheumatic on other grounds.

Course and prognosis The average life expectancy of rheumatic aortic incompetence is 20 to 30 years from its development. Prognosis should be based on the size of the left ventricle and upon the degree of incompetence as judged by peripheral vascular behaviour. Effort tolerance often remains remarkably good until near the end. Failure is commonly with normal rhythm and is usually left ventricular at first. Complications are practically limited to bacterial endocarditis.

4 AORTIC STENOSIS

Pathology Fibrous scar tissue representing healed aortic valvulitis usually causes fusion of the cusps at their commissures. Slight narrowing at the aortic aperture is thus found in most cases of rheumatic aortic valve disease. When fusion extends further up the margins of the cusps true stenosis results. Valve leaflets become thick, rigid, distorted and often unrecognisable. Secondary valve calcification is common. The aorta and large arteries often remain remarkably free from atheroma.

Effect on function The aortic orifice must be reduced to about one quarter of its natural size before changes in the circulation can be demonstrated (Wiggers 1935). Left ventricular pressure curves then show a raised initial tension, steep isometric pressure gradient and an elevated maximum pressure that is reached relatively early in systole but there is no collapse as in aortic incompetence. Pressure curves obtained from the aorta show an initial steep rise interrupted by an anacrotic notch and followed by a slower rise that reaches its maximum late in systole; the maximum pressure attained is less than normal. The more severe the stenosis the earlier the anacrotic notch. The ejection phase is prolonged.

To maintain the stroke volume and cardiac output great power must be developed by the left ventricle. The chamber is more hypertrophied and less dilated than in aortic incompetence. Again the left ventricle must have sufficient time to perform its task; it needs a long stroke and requires to be well filled in diastole.

Clinical features Aortic stenosis is at least twice as common in men as in women. The lesion may be discovered at any time from adolescence to old age, usually in the sixth decade. Female patients tend to be younger than male.

Patients with aortic stenosis may complain of syncope (10 per cent) of angina pectoris (20 per cent) or of symptoms referable to left ventricular or congestive heart failure (Contratto and Levine 1937). Syncope is of two kinds, cardiac and vasomotor. Cardiac syncope is abrupt and fleeting and

may be due to paroxysmal ventricular fibrillation or possibly to locking of the valve (de Veer 1938). Such attacks herald sudden death from a similar mechanism. The low blood pressure of aortic stenosis predisposes to vasomotor and to orthostatic syncope. Angina pectoris appears to depend upon poor coronary filling secondary to a jet effect or to the low mean aortic pressure. Attacks are in no way different from those due to coronary atherosclerosis. As usual, women with angina tend to be a decade older than the men.

Fig. 9 21—Arteriogram in a case of aortic stenosis. The percussion wave is prolonged and the maximum pressure is reached late in systole.

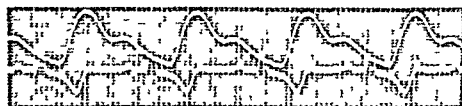


Fig. 9 21—Arteriogram in a case of aortic stenosis. The percussion wave is prolonged and the maximum pressure is reached late in systole.
(Bourne, J. D., F. H. Gardner and M. Z. B.)

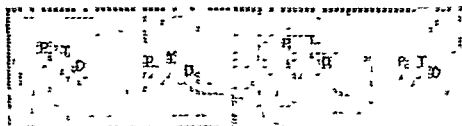


Fig. 9 22—Arteriogram illustrating pulsus bisferiens in a case of combined aortic stenosis and incompetence. P is the percussion wave, T the tidal wave, both are systolic events.
(Bourne, J. D., F. H. Gardner and M. Z. B.)

The physical signs are as follows:

1. There is often a delicate pale pink complexion—the Dresden china look.

2. The pulse is characteristic when relatively slow (fig. 9 21), being small and sustained (plateau or slow rising pulse). It depends upon the longer duration of left ventricular systole, the low blood pressure, and upon the delayed development of maximum aortic pressure. These features tend to disappear as the heart rate quickens. When aortic incompetence is present as well, the pulse assumes a bisferiens quality (fig. 9 22). To the palpating finger it feels double and may even be mistaken for coupling.

due to premature ectopic beats. Both waves occur during the ejection phase. According to Bramwell (1937) the second impulse is tidal in nature, being due to overlapping and partial fusion between a forcible but prolonged percussion wave and its reflection from the periphery. Aortic incompetence increases the force of the percussion wave, aortic stenosis prolongs it. Neither alone will produce this pulse.

3. The blood pressure is variable. In severe cases it is low, and the pulse pressure is small, but in mild or moderate cases, or when there is recognizable aortic incompetence, it may be elevated and the pulse pressure may be increased. About 10 per cent are truly hypertensive—an incidence a good deal lower than in controls of the same age group.

4. The apex beat is displaced downwards and to the left, and the cardiac impulse is slow and heaving. The left ventricle is hypertrophied rather than dilated.

5. A basal systolic thrill is usually present. It is best appreciated when the patient leans forward and stops breathing in full expiration. It may be most intense either to the right or to the left of the sternum. A systolic thrill may also be felt over the carotid or subclavian arteries. Although such a thrill is not diagnostic of aortic

stenosis, it is suggestive and encourages prolonged search at the base.

6. A long, rough, basal systolic murmur is almost invariable. It is conducted into the cervical arteries and may sometimes be heard remarkably well at the apex beat. The second heart sound is usually soft or absent.

7. On fluoroscopy the left ventricle looks dense and bulky. The aorta may be conspicuous or relatively hypoplastic (fig. 9.23). Calcification of the aortic valve can be seen in most cases, particularly if the patient is over 50.

8. The electrocardiogram usually provides convincing evidence of left ventricular enlargement. Perhaps owing to the concentric type of hypertrophy and to the lack of dilatation, the heart is often electrically vertical. Standard leads then show the concordant pattern of left ventricular preponderance (fig. 9.24). Exceptionally high voltage R waves are characteristic of aortic stenosis. T is frequently inverted in leads facing the surface of



Fig. 9.23—Skilogram of a case of aortic stenosis showing great enlargement of the left ventricle, slight prominence of the ascending aorta, and nilar congestion.

the left ventricle. Left bundle branch block occurs in about 15 per cent of cases, auricular fibrillation in about 5 per cent.

Differential diagnosis. If as much attention were paid to the quality of the peripheral pulse as to cardiac murmurs, aortic stenosis would be both less frequently overlooked and less often diagnosed in error. Functional basal systolic murmurs should not cause confusion. The murmur and thrill

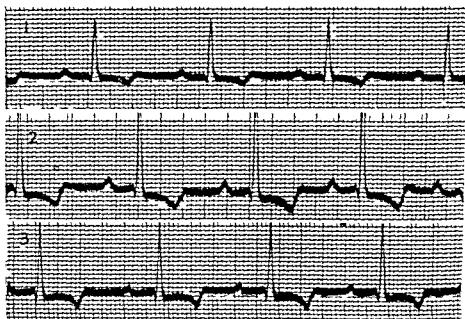


Fig 9-24—Electrocardiogram in a case of aortic stenosis showing concordant left ventricular preponderance in standard leads, the heart being vertical.

of ventricular septal defect though lower and more to the left may be more difficult to distinguish but the normal peripheral pulse, normal electrocardiogram, X-ray appearances and absence of valve calcification help to prevent mistakes. Organic mitral incompetence may be recognised by the normal peripheral pulse and by the size and behaviour of the left auricle when viewed fluoroscopically. The site of the maximum intensity of the murmur is less reliable evidence.

Etiological diagnosis is more difficult. Rheumatic aortic stenosis must be distinguished from congenital and calcific atherosclerotic varieties. In congenital sub-aortic stenosis symptoms are absent, the heart is little enlarged, peripheral vascular findings are minimal, there is no aortic incompetence and calcification is rare. Congenital valvular stenosis may be indistinguishable from the rheumatic variety but the lesion is usually discovered in childhood, growth is retarded, the aorta may be hypoplastic and incompetence does not occur.

It is uncertain whether calcific aortic stenosis in elderly or middle aged

subjects is atherosclerotic or rheumatic. Thus eleven of twenty one cases reported by Christian (1931) gave a history of rheumatic fever. Dry and Willis (1939) obtained a rheumatic history in 22 per cent of 228 cases and Clawson, Noble and Iufkin (1938) found a rheumatic history in 35 per cent of 200 cases. On the other hand in the quoted series of Dry and Willis there were 91 necropsied cases without disease of other valves; a rheumatic history was obtained in only four of these—the usual incidence in any series of normal controls. Again in the quoted series of Clawson and his colleagues 20.5 per cent of the patients were under 41 years of age and 39 per cent were under 51; moreover 89 had a mitral lesion as well. It is obvious that many of these cases were rheumatic but this has little bearing upon the question of whether or not pure calcific aortic stenosis in elderly people is rheumatic. On the pathological side Clawson (1931) particularly has drawn attention to the frequency of inflammatory stigmata of the rheumatic type but others notably Soval and Gross (1936) have been unable to confirm such findings. The best evidence of a rheumatic or other inflammatory etiology is perhaps the remarkable absence of atherosclerosis in the aorta and coronary arteries in most cases. All observers have agreed on this point that these vessels must have been long protected by the stenosis. However Monckeberg's original thesis that calcific aortic stenosis in elderly subjects may be degenerative (Monckeberg 1904) has not been altogether disproved.

Clinically calcific aortic stenosis in elderly subjects behaves like rheumatic aortic stenosis.

Course and prognosis. At least 15 per cent of aortic stenotic subjects die abruptly particularly if they have suffered from cardiac syncope or from angina pectoris. About 10 per cent develop bacterial endocarditis. The majority who survive these risks succumb to congestive heart failure sooner or later.

The prognosis should be based upon the behaviour of the peripheral pulse upon the size of the left ventricle and upon the nature of the symptoms. When the pulse and left ventricle are relatively normal the outlook is good and life expectancy little curtailed. Cardiac syncope and angina pectoris are serious and give warning of sudden death at any time. Between these two extremes all grades of severity are encountered. Most cases however enjoy good effort tolerance until well into middle age.

5 TRICUSPID INCOMPETENCE

Tricuspid incompetence may be functional or organic the former being secondary to right ventricular dilatation with expansion of the tricuspid ring. Clinical distinction is difficult in the first instance but the course and response to digitalis and to rest may clarify the issue. Functional incompetence may be temporary organic tricuspid disease is always permanent.

The diagnosis is based upon the following features

- 1 The cervical veins are engorged and pulsate with extraordinary

vigour. The quality of venous pulsation is altered when there is auricular fibrillation which is usual, a single prolonged venous pulse may replace the normal double movement. The *c* and *v* waves of the jugular phlebogram are more or less fused (fig 9 25). The normal drop in venous pressure that follows the *c* wave is due to systolic descent of the base (floor of the auricles or atrio-ventricular septum) which produces a sucking effect. In tricuspid incompetence this negative pressure is replaced by partial transmission of the right ventricular systolic pressure to the right auricle (Bloomfield

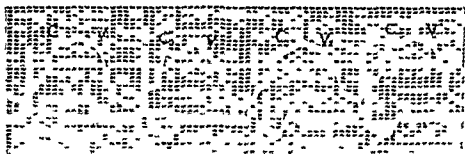


FIG 9 25—Jugular phlebogram showing fusion of the *c* and *v* waves in a case of tricuspid incompetence. Owing to auricular fibrillation the *a* wave is absent.

(B. L. S. J. D. M. Zorab)

et al. 1946). When there is normal rhythm the ventricular form of venous pulse may be preceded by a powerful *a* wave; the jugular pulse is then double, but the abnormal form may still be recognised at the bedside.

2. In long-standing cases a brownish pigmentation may be seen in the skin, especially of the head and neck.

3. Systolic expansile pulsation of a considerably enlarged liver can usually be recognised by palpation. This must be distinguished from transmitted right ventricular pulsation.

4. There is usually an early, long, blowing systolic murmur low down the left border of the sternum, and there may be an associated thrill. The murmur may increase in full inspiration.

5. X-rays show gross dilatation of the right auricle, the border of which meets the diaphragm at a right angle, or even obtusely (fig 9 26). In pericardial effusion this angle is usually acute. On fluoroscopy the right auricle occasionally expands in systole, and the right lobe of the diaphragm may reflect hepatic pulsation.

6. Catheter studies have demonstrated reversal of the central venous pressure gradient during systole, forward flow being limited to diastole (Bloomfield *et al.* 1946). Venous valves take on the function of the tricuspid valve. The diagnosis of tricuspid incompetence may thus be proved by demonstrating a higher mean pressure in the right auricle and superior vena cava than in the subclavian vein (Sharpey-Schafer 1947).



Fig 926—Sk gram showing gross dilatation of the right auricle with a blunt right cardio phrenic angle in a case of tricuspid incompetence

moreover, as the catheter is withdrawn pulsation ceases abruptly the moment the pressure falls (fig 9 27)

The recognition of organic or permanent tricuspid incompetence is important, because patients so afflicted would otherwise be kept in bed indefinitely in the belief that they suffered from congestive heart failure. Yet these patients may remain remarkably free from symptoms and may be able to carry on their daily work for years despite gross physical signs.

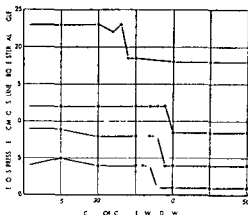


Fig 9 27—Graph illustrating fall in mean central venous pressure as the catheter is withdrawn from right auricle and superior vena cava into the subclavian vein

accompanied by mitral stenosis (Pitt 1909) often by aortic valve disease as well.

Tricuspid stenosis prevents proper cardiac filling and in this respect resembles constrictive pericarditis (Thompson and Levine 1937). Pulmonary congestion threatened by other valve lesions is thus prevented; attacks of dyspnoea and orthopnoea are noticeably absent; systemic venous engorgement, hepatic distension, ascites and oedema occur instead. Tricuspid incompetence is usually associated.

The diagnosis is based upon the following findings:

1. Engorgement of the cervical veins. The characteristic finding is an abrupt and powerful *a* wave which may be called a venous Corrigan; this does not alter with change of posture. It is of course only present in cases with normal rhythm.

2. Brownish discolouration of the skin, especially of the head and neck as in tricuspid incompetence. Jaundice may also occur.

3. Considerable enlargement of the liver, sometimes progressing to cardiac cirrhosis. The organ is unduly firm and is not tender. Pulsation is presystolic when due to transmission of a giant *a* wave or systolic if there is associated tricuspid incompetence. Ascites is common.

4. A tricuspid diastolic murmur heard best low down the left border of the sternum, similar in quality and timing to the mitral diastolic murmur. It may be accompanied by a thrill. As mitral stenosis is nearly always present, a separate tricuspid murmur easily escapes notice.

6 TRICUSPID STENOSIS

Although organic disease of the tricuspid valve is found at necropsy in 10 to 15 per cent of all cases of chronic rheumatic heart disease (Smith and Levine 1942), clinical tricuspid stenosis is infrequently recognised. It is nearly always

5 Gross dilatation of the right auricle seen on fluoroscopy

6 Tall sharp P waves in the electrocardiogram in cases with normal rhythm (fig 9 28) They represent right auricular hypertrophy Owing to coincident mitral stenosis the P waves are usually widened as well Atrial fibrillation however is often present

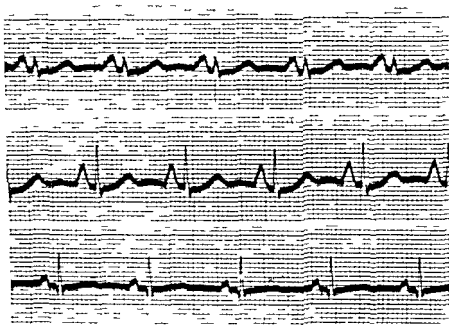


Fig 9 28—Electrocardiogram showing exceptionally tall P waves in a case of mitral and tricuspid stenosis

The most significant fact about tricuspid disease is that it gives rise to gross signs suggesting severe right ventricular failure in patients who are up and about and who may be practically free from symptoms the discrepancy should at once draw attention to the diagnosis Life expectancy averages about five years from the time the diagnosis is first made (Aceves and Carral 1947)

MYOCARDIAL FIBROSIS

The rheumatic process affects the heart muscle as well as the valves leaving patchy myocardial fibrosis Occasionally cases of heart failure come to necropsy in which nothing but patchy fibrosis is found It is possible that some of these represent old rheumatic carditis Again failure may occur in rheumatic heart disease when valvular scarring is insignificant Permanent heart block or bundle branch block may be caused by fibrosis at the appropriate site

ADHERENT PERICARDIUM

Pericarditis is one of the more innocent rheumatic lesions in respect to its after effects. Sometimes the two layers of the sac are fused and thickened but this causes no trouble. Secondary calcification is rare and scanty when present. Occasionally adhesions form between the pericardium and surrounding structures so that the heart becomes anchored firmly in the mediastinum or to the thoracic wall. The apex beat does not shift with change of posture and Broadbent's sign is positive. Current opinion favours the view that this too is relatively innocent. Pick's disease or chronic constrictive pericarditis is very rarely if ever rheumatic. The matter is more fully discussed in Chapter XII.

TREATMENT

Competitive effort and hard physical work should be forbidden. Precautions should be taken against exposure to cold and infection. Patients are advised to train for a sedentary occupation as a safeguard against progressive cardiac enlargement and future breakdown. In mild cases moderate physical exertion may be allowed but the patient must live well within the limits of effort tolerance.

No drugs, no particular diet and no special measures are required except for complications which must be treated as they arise as described elsewhere. Intercurrent pyogenic infections call for prompt chemotherapy. The problem of pregnancy is discussed elsewhere (page 507).

Modern surgery offers some hope for patients with tight mitral stenosis and severe pulmonary congestion. The most promising operation is mitral valvulotomy as performed by R. C. Brock (1950). Anastomosing a pulmonary vein to the azygos vein (D. Allaines *et al.* 1949) certainly relieves congestion in the lesser circulation but presumably lowers the cardiac output and is unlikely to be so effective ultimately. The same may be said for the more hazardous procedure of tricuspid valvulotomy.

REFERENCES

- Aceves S. and Carral R. (1947) The diagnosis of tricuspid valve disease. *Amer Heart J* 34 114.
- Baker C. Brock R. C. and Campbell M. (1950) Valvulotomy for mitral stenosis. *BMJ* 1 1283.
- Berconsky I. and Neuman J. (1945) Frecuencia y significado de la Hipertension arterial en la estrechez mitral. *Rev. Argent Cardiol* 12 94.
- Bloomfield R. A. Lauson H. D. Cournand A. Breed E. S. and Richards D. W. (1946) Recording of right heart pressures in normal subjects and in patients with chronic pulmonary disease and various types of cardiocirculatory disease. *J clin Invest* 25 639.
- Boone J. A. and Levine S. A. (1938) The prognosis in potential rheumatic heart disease and rheumatic mitral insufficiency. *Amer J med Sc* 195 764.
- Bramwell C. (1937) Arterial pulse in health and disease. *Lancet* ii 239 301.

Braun Menendez E and Orias O (1935) Curacion de las fases del ciclo cardiaco en hipertensos *Rev Argent Cardiol* 2 186

Brock R C (1950) See Baker

Cabot R C (1926) Facts on the heart Philadelphia

de la Chapelle C E Graef I and Rottino A (1934) Studies in rheumatic heart disease analysis of 119 hearts with special reference to relationship of auricular fibrillation to mitral valvular deformity and certain rheumatic tissue changes *Amer Heart J* 10 6

Christian H A (1931) Aortic stenosis with calcification *J Amer med Ass* 97 158

Clawson B J (1931) Nonsyphilitic aortic valve deformity *Arch Path* 12 889 — Noble J F and Lufkin N H (1938) "The calcified nodular deformity of the aortic valve" *Amer Heart J* 15 58

Contratto A W and Levine S A (1937) Aortic stenosis with special reference to angina pectoris and syncope *Ann intern Med* 10 1636

Coombs C F (1924) Rheumatic heart disease Bristol

Corrigan D J (1832) On permanent patency of the mouth of the aorta or inadequacy of the aortic valves *Edin med and surg J* 37 225

D Allaines F Lenègre J Dubost Ch Mathivat A and Seebat L (1949) Retrecissement mitral Anastomose veine pulmonaire veine azygos Premier cas opere *Mémoires de l'Académie de Chirurgie Paris* 75 318

Davis D and Weiss S (1931) Rheumatic heart disease I Incidence and role in the causation of death A study of 5,215 consecutive necropsies *Amer Heart J* 7 146

Dry T J and Willius F A (1939) Calcareous disease of the aortic valve *Ibid* 17 138

Durozier P (1861) Du double souffle intermittent crural comme signe de l'insuffisance aortique *Arch gen de Med Paris* 107 417 588

Evans W (1947) Heart murmurs *Brit Heart J* 9 1

Fauvel S A (1843) Memoire sur les signes stethoscopiques du retrecissement de l'orifice auriculo ventriculaire gauche du coeur *Arch gen de Med Paris* (ser 4) 1 1

Fetterolf G and Norris G W (1911) The anatomical explanation of the paralysis of the left recurrent laryngeal nerve found in certain cases of mitral stenosis *Amer J med Sc* 141 625

Flint A (1862) On cardiac murmurs *Ibid* 44 29

Gardner W T (1861) A short account of cardiac murmurs *Edin Med J* 7 445

Gouley B A (1938) The role of mitral stenosis and of post rheumatic pulmonary fibrosis in the evolution of chronic rheumatic heart disease *Amer J med Sc* 196 11

de Graff A C and Lingg C (1935) Course of rheumatic heart disease in adults influence of auricular fibrillation on course of rheumatic heart disease *Amer Heart J* 10 459

Gumpert T E (1947) Miliary appearances in the lungs in mitral stenosis *Brit med J* n 488

Harrison T R (1935) Failure of the circulation Baltimore

Hope J (1839) A treatise on the diseases of the heart and great vessels 3rd ed London

King T W (1838) On morbid flattening or compression of the left bronchus produced by dilatation of the left auricle *Guy's Hosp Rep* 175

Kuttner A G and Markowitz M (1948) The diagnosis of mitral insufficiency in rheumatic children *Amer Heart J* 35 718

Luisada A A (1943) On the pathogenesis of the signs of Traube and Durozier in aortic insufficiency A graphic study *Ibid* 26 721

Margolies A and Wolferth C C (1932) The opening snap (Claquement d'ouverture de la mitrale) in mitral stenosis its characteristics mechanism of production and diagnostic importance *Ibid* 7 443

Mönckberg J G (1904) Der normale histologische Bau und die Sclerose der Aortenklappen *Virchows Arch f path Anat* 176 472

Ortiz y Ramirez (1933) Una Nueva Teoria de los soplos anorganicos frotamientos cardio serosos *Arch Lat Am d Cardiol y Hematol* 3 45

Ortner N (1897) Recurrenslähmung bei mitralstenose *Wien klin Wschr* 10 753

Parkinson J (1945) Rheumatic fever and heart disease *Lancet* ii 657 —

and Harley R (1946) Early diagnosis of rheumatic valvular disease in recruits *Brit Heart J* 8 212

Pitt G N (1909) The system of medicine ed Allbutt and Rolleston London 6 330 — (1909) Right sided valvular diseases *Syst Med Allbut and Rolleston* London 7 310

Rolleston H (1940) History of aortic regurgitation *Ann Med Hist* 2 271

— (1941) The history of mitral stenosis *Brit Heart J* 3 1

Sharpey Schafer F P (1947) Unpublished observations

Smith J A and Levine S A (1942) The clinical features of tricuspid stenosis *Amer Heart J* 23 739

Soval A R and Gross L (1936) Calcific sclerosis of the aortic valve *Arch Path* 22 477

Steel G (1888) The murmur of high pressure in the pulmonary artery *Med Chronicle Manchester* 9 18.

Thompson W P and Levine S A (1937) Note on duration of symptoms and age at death in chronic rheumatic valvular disease especially in tricuspid stenosis *Amer J med Sc* 193 4

de Veer J A (1938) Sudden death in aortic stenosis explanation on a mechanical basis *Amer Heart J* 15 243

Watson T (1843) Principles and practice of physic London

Werner S C (1936) Rheumatic cardiac disease Association of active rheumatic fever with heart failure *Arch intern Med* 57 94

Wiggers C J (1928) Pressure pulses in the cardiovascular system London — (1935) Physiology in health and disease London

CHAPTER X

OTHER FORMS OF CARDITIS

THE HEART IN DIPHThERIA

DIPHThERIA may cause peripheral circulatory collapse or toxic myocarditis. Cutaneous diphtheria so easily overlooked and so often untreated until too late may be as lethal as the common faucial type. Early and adequate treatment with antitoxin has greatly reduced the incidence of toxic complications but has by no means abolished them.

CIRCULATORY COLLAPSE

Towards the end of the first week or during the second week of the illness the blood pressure may fall well below 100 mm Hg the patient becomes faint sick and restless the skin pale cold and clammy the pulse rapid and thready. Loss of vasomotor tone may be due to toxic depression of the vasomotor centre perhaps to peripheral sympathetic paresis or possibly to poisoning of the vessels themselves. Occasionally it is brought about by suprarenal failure due to necrosis or hæmorrhage. The earlier the onset of circulatory collapse the worse the prognosis. Patients usually remain in a critical state for several days in those who recover improvement may then occur but the blood pressure usually remains low for two or three weeks.

The course of diphtheria may be complicated (as well as alleviated) by serum therapy for this may induce not only immediate collapse from anaphylactic shock in a sensitised individual but also later collapse from loss of plasma into the tissue spaces associated with serum sickness. Urticaria and œdema usually on the ninth day may be extreme and result in a diminished blood volume and hæmoconcentration. Diphtheritic circulatory collapse and allergic shock may thus be expected at about the same time and diagnostic difficulties may arise.

Treatment of both conditions consists of absolute rest adrenaline 7 to 10 minims (0.4 to 0.6 ml) subcutaneously two to four hourly and of raising the foot of the bed. Sodium salicylate 20 grains (1.3 G) two hourly and the antihistamine drugs are helpful in serum sickness. The limbs may be bandaged with advantage in diphtheritic circulatory failure and the cautious infusion of plasma is not irrational. The prognosis is grave.

TOXIC MYOCARDITIS

Pathology. Diphtheritic carditis being toxic in nature may prove fatal without causing advanced changes in morbid histology. The characteristic

finding is hyaline degeneration or necrosis of muscle the fibres losing their striations and presenting a swollen granular appearance. Lesions are patchily distributed and only short segments of individual muscle fibres may be affected. Monocytes cluster round the debris and fibroblastic repair follows.

Clinical features Disturbances of rhythm tend to occur first usually during the second week of the disease. Partial or complete heart block and bundle branch block are the best known and in patients who recover from the illness are usually but not invariably transient (Peiry 1939). Both heart block and bundle branch block commonly denote severe carditis most such cases proving fatal (Burkhardt, Iggleson and Smith 1938). Ectopic beats are common and although often innocent and unrelated to carditis should be viewed with suspicion in diphtheria. Auricular fibrillation and paroxysmal tachycardia are rare. Ventricular fibrillation may be responsible for sudden death.

Other evidence of carditis tends to occur a little later usually during the third week. Sinus tachycardia, gallop rhythm, enlargement of the heart and reduction of the pulse pressure are usual. The onset of heart failure may be suggested by pallor, breathlessness, præcordial oppression and vomiting. Congestion is systemic rather than pulmonary the jugular venous pressure being raised and the liver distended; there is rarely orthopnoea, paroxysmal cardiac dyspnoea or pulmonary oedema. Significant murmurs and pericardial friction are absent.

The electrocardiogram is especially helpful in the diagnosis of diphtheritic carditis much more so than in rheumatic carditis. Depression of the RS-T segment or primary inversion of the T wave in most leads is characteristic and is found during the second week in the majority of cases which develop clinical carditis and in some that do not. A similar pattern may be produced in cats within 48 hours by injecting diphtheritic toxin (Nathanson 1928). Of 600 cases of diphtheria studied by Altshuler *et al* (1948) 108 or 18 per cent developed these changes while only 11 showed heart block.

Radiological studies on diphtheritic carditis are rare because patients are not allowed to stand or sit and should not be moved to the X-ray department. Portable skiagrams give little information about the size of the heart.

Prognosis The outlook is grave for sudden death is common and presumably results from ventricular fibrillation or asystole. Some patients die from congestive heart failure. Not infrequently associated circulatory failure complicates the picture. Those who survive usually develop polyneuritis later and this is apt to be severe. The total mortality rate is difficult to assess for mild cases may well be overlooked but it is usually put at 50 per cent.

If the patient survives the ultimate prognosis is excellent (White *et al* 1937) and complete recovery may be promised without reserve. It is

important that the patient should be convinced of this from the start in order to prevent anxiety neurosis and to maintain good morale

Treatment Antitoxic serum will already have been administered in most cases if not it is too late to give it by the time cardiovascular symptoms develop. The axiom that antitoxin cannot do any harm and might as well be given even at this stage is untrue for serum reactions are common and may prove fatal when there is toxic circulatory collapse or carditis

Prophylactic treatment in addition to early and adequate doses of antitoxin consists of complete rest in bed for a minimum period of one month in all cases of diphtheria. If by the end of this time there is no evidence of cardiovascular or neuro intoxication there is little further risk to life. Should any such intoxication have occurred however bed rest must be extended for another month otherwise sudden death may occur during convalescence in the second month. Patients may be treated with far less respect subsequently even when they have extensive polyneuritis

The treatment of recognised carditis is unsatisfactory. Absolute rest is essential for sudden slight effort even sitting up in bed may prove fatal during the critical period. Patients should be nursed flat with one pillow and should have everything done for them including being fed and washed

Diet should be light and fluids limited to two pints daily. Digitalis is dangerous and should only be used in rare cases when auricular fibrillation with a rapid ventricular rate is associated with severe congestive heart failure. Auricular fibrillation without failure is unlikely to last long if untreated and is less dangerous than digitalis. Diphtheritic heart failure with normal rhythm responds poorly to digitalis and the drug is usually better withheld

THE HEART IN ACUTE INFECTIONS

Up to the beginning of the twentieth century it was generally believed that toxic carditis was a common complication of certain fevers such as influenza. It came to be recognised however that although cloudy swelling and fatty degeneration were often found at autopsy in cases dying from severe general infections clinical evidence of cardiac involvement was rare. The change of view followed the establishment of stricter criteria for diagnosing organic heart disease. Palpitations and irregularities of the heart were shown to be due to autonomic disturbance or to innocent ectopic beats. Systolic murmurs lost their previous significance. Effort syndrome following infections was proved attributable to anxiety. X rays failed to confirm clinical cardiac enlargement (based on the position of the apex beat). Standard lead electrocardiograms were rarely abnormal. The weight of negative evidence was considerable and it became the custom to recognise no form of carditis other than that due to rheumatism or diphtheria. In recent years however the earlier view has gained some support

Thus Burch and Reaser (1947) considered infective or toxic carditis to be the most common cause of organic heart disease. Gore and Saphir (1947) found that diphtheria and rheumatism accounted for less than 25 per cent of fatal cases of myocarditis; they contended that carditis was common in a host of infectious and protozoal diseases, intoxications and allergic states including especially scrub typhus, bacterial endocarditis, meningococcal septicæmia and sulphonamide allergy. At the same time an increasing number of cases of isolated myocarditis (Fiedler's) have been reported. It may be as well therefore to review the known facts critically for there is grave danger that this modern swing back may go too far.

FAILURE OF THE PERIPHERAL CIRCULATION

Cardiovascular disturbances in acute infections are commonly of two kinds and neither is due to a cardiac fault. The first is peripheral circulatory failure. This may be due to depression of the vasomotor centre, to toxic paresis of the vessels themselves, to suprarenal failure or to diminution of the blood volume from dehydration or from loss of plasma into the tissue spaces through damaged vessels. The essential mechanism is critical discrepancy between the effective vascular capacity and the blood volume and the chief clinical feature is low blood pressure.

A good sign of vascular relaxation is a markedly dirotic pulse and although not necessarily serious should put the physician on guard. Another significant feature is pallor and coldness of the extremities due to vasoconstriction in the skin; this appears to be a compensatory mechanism helping to maintain the venous pressure and blood pressure when dangerous vasodilatation occurs elsewhere, e.g. in muscle. Impending failure of compensatory vasoconstriction may be indicated by waxing and waning of the systolic blood pressure through a range of 10 to 20 mm. Hg. A fourth indication of circulatory failure is mental confusion or faintness in the sitting posture. Whilst tachycardia is the rule and the half hourly pulse chart of some value, it should be understood that deceleration sometimes accompanies a falling blood pressure and that the character of the pulse is as important as its rate.

Circulatory failure should be treated by nursing the patient flat or with the foot of the bed raised and by the intravenous administration of serum or plasma by the drip method with or without 0.5 to 1 mg. of adrenaline to the bottle.

The second common cardiovascular reaction to acute fevers is vasomotor neurosis during convalescence. This is discussed in Chapter XVI.

TOXIC MYOCARDITIS

True toxic myocarditis does occur, however, especially perhaps in pneumonia. Sections reveal focal hyaline necrosis, i.e. granular degeneration and loss of striation of the muscle fibres, patchily distributed. Cellular reaction with monocytes predominating and fibroblastic repair follow—as in diph-

theritic carditis which it resembles. This histological picture is common to most forms of carditis—hence the difficulty in making an etiological diagnosis from autopsy findings. For example, 35 cases of sudden death following tonsillitis or common cold were reported by Gore and Saphir (1947) and ascribed to toxic myocarditis. Thirty one of them, however, could have been due to diphtheria or pneumonia; a negative throat swab does not exclude diphtheria.

Myocarditis and diffuse glomerulonephritis have long been known to complicate bacterial endocarditis, but when the death rate of the septicæmic stage was 98 per cent, they received relatively little attention. Since the introduction of penicillin, however, heart failure from myocarditis has become chiefly responsible for the present 25 per cent mortality.

Histological examination of the heart in cases dying from meningococcal infection may disclose evidence of carditis, but clinical signs of cardiac involvement are most unusual, and the total mortality rate in adults is less than 1 per cent (Daniels *et al.* 1943). Whether meningitic or septicæmic in form, the infection responds particularly well to sulphonamides, and the rare occurrence of clinical carditis would probably be attributed to sulphonamide allergy.

Allergic forms of carditis, sometimes with peri arteritis, giant cells, eosinophils, and nodules, have been described (Reinhart 1946). Sulphonamides provide an example of an antigen which may provoke such a response (French and Weller 1942; French 1946). Carditis, pericarditis, and endocarditis may accompany acute disseminated lupus (Humphreys 1948).

A most convincing type of protozoal carditis may accompany South American trypanosomiasis or Chagas disease (Chagas 1909). Leishmanial forms of *T. cruzi* multiply chiefly in the cells of the heart, brain, and liver; the affected cells finally rupture and liberate the parasites into the blood stream. An intense local inflammatory reaction follows. The signs and symptoms of a typical acute or subacute carditis may dominate the clinical picture, and sudden death is common (Mosely and Miller 1945).

Carditis accompanying scrub typhus (Tsutsugamushi fever) is less convincing. Although histology may reveal myocardial damage and cellular infiltration in fatal cases (Corbett 1943), the clinical course of the disease seems to be little influenced by them (Williams *et al.* 1944; Berry *et al.* 1945). In a series of 184 cases seen within one to four weeks after the acute symptoms had subsided, and 10 cases seen during the stage of fever, the electrocardiogram was virtually normal (Howell 1945). For further information, the reader is referred to the issue of the *American Journal of Hygiene*, May 1945, which is devoted to studies on scrub typhus.

To assess the clinical value of the work of Gore and Saphir quoted above, it is worth noting that 16 per cent of their 1,402 cases of myocarditis were due to scrub typhus, and there was no evidence that myocarditis was the cause of death. Their cases were highly selected, excluded children, and

were based entirely on autopsy findings there were 227 examples of scrub typhus 208 of bacterial endocarditis 144 of diphtheria 130 of rheumatic carditis and 105 of sulphonamide allergy The reader will draw his own conclusions

Clinically significant carditis accompanying acute infections in Great Britain (other than rheumatic fever diphtheria and bacterial endocarditis) is undoubtedly rare This applies as much to typhoid (Porter and Bloom 1935) and influenza (Wood 1941) as to the common exanthemata

Clinical features of toxic myocarditis In acute cases the signs and symptoms are similar to those of diphtheritic myocarditis except that they may

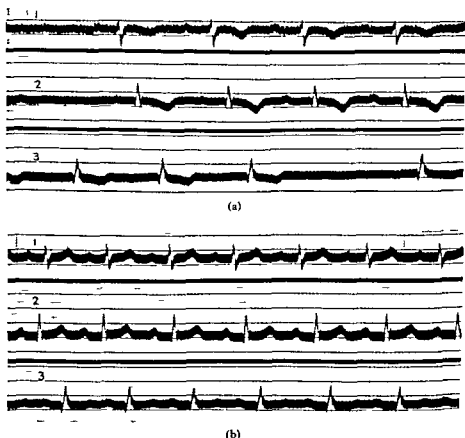


Fig 1001—Electrocardiogram in a case of toxic myocarditis due to pneumonia
(a) Shows partial heart block with dropped beats and inversion of the T wave in all leads
(b) After recovery

occur earlier during the febrile stage of the infection Symptoms attributable to cardiac involvement may be absent on the other hand there may be dyspnoea unexpected vomiting pallor and peripheral cyanosis due to congestive failure substernal oppression or discomfort or palpitations

associated with changes of rhythm. It may be difficult to distinguish cardiac symptoms from those due to general toxæmia particularly when there is peripheral circulatory failure. Sudden death is not infrequently the first tragic proof of myocarditis.

Physical signs include a small rapid thready pulse, low systolic blood pressure, small pulse pressure, gallop rhythm, dilatation of the heart, congestive heart failure, abnormalities of rhythm and electrocardiographic changes. The small rapid pulse and the low blood pressure may equally well be due to peripheral circulatory failure and the gallop rhythm to fever (especially when there is anæmia). The size of the heart may be difficult to assess under the clinical circumstances and the patient should not be moved to the X-ray department for more exact information. The importance of recognising early signs of congestive heart failure will thus be appreciated. Abnormalities of rhythm are also important and include all grades of heart block, auricular flutter or fibrillation and paroxysmal tachycardia. The electrocardiogram is especially helpful not only in establishing the nature of a rhythm change but also in revealing partial heart block and abnormalities of the T wave (fig. 10.01).

Sometimes the course of toxic myocarditis is subacute or chronic. The clinical features then closely resemble those of isolated (Fiedler's) myocarditis described on page 318.

Prognosis. If the diagnosis is beyond doubt the outlook is grave, the mortality rate probably approaching 50 per cent. Whether central or peripheral in mechanism, the combination of hypotension and a small rapid pulse is always dangerous and congestive heart failure often proves fatal. Abnormalities of rhythm and alterations of the T wave without the manifestations just mentioned are perhaps less serious.

Many cases of mild toxic myocarditis must pass unrecognised but this is not a matter for concern for recovery appears to be complete in all non-fatal cases.

Treatment. Bed rest and specific chemotherapy (when applicable) for all acute infections are axiomatic; bed rest should be absolute if the cardiovascular system is involved. The patient should be nursed in the position of maximum comfort but if the blood pressure is below 100 mm. Hg and there is no evidence of congestive failure he should be kept horizontal; if there is congestive failure he should be propped up at 30 to 45 degrees against a back rest. Digitalis should be avoided unless there is frank congestive failure for it increases the risk of sudden death from ventricular fibrillation and may aggravate minor degrees of heart block. If the venous pressure is well raised and the liver distended however it should not be withheld and it may be invaluable in cases of auricular flutter or fibrillation. Mersalyl and a low sodium diet may be given if there is fluid retention. Quinidine 3 to 5 grains (0.2 to 0.3 G.) t.d.s. may prevent paroxysmal rhythm changes including ventricular fibrillation.

It must be admitted however that toxic myocarditis is little influenced

by therapy and is apt to be fatal or otherwise according to its severity and irrespective of treatment. As in diphtheria nearly all who recover do so completely.

ISOLATED MYOCARDITIS

Isolated myocarditis (Scott and Saphir 1929) is a subacute inflammation of the heart of unknown etiology characterised by patchy myocardial necrosis, cellular infiltration and fibroblastic repair as in other forms of myocarditis. It was first properly described by Friedler (1899). The disease may not be a specific entity and is difficult to distinguish pathologically from known forms of toxic or infective myocarditis of relatively long duration.

Incidence. Although still relatively rare isolated myocarditis is being recognised with increasing frequency. The majority of cases have occurred in subjects between the ages of 20 and 50 but infants, children and old people are not exempt. The disease has been reported sporadically in most countries and races; the only possible minor epidemic occurred in African troops serving in the Middle East (Bedford and Konstam, 1946), but neither the nature of the carditis nor its infective origin were certain; there were 40 cases with 17 deaths. They may well have had a basis of malnutrition.

Pathology. Patchy necrosis of muscle is thought to be the primary lesion (fig. 10.02). Cellular reaction may be focal or more diffusely interstitial. Monocytes predominate but in the acute stage polymorphs may be more numerous. Haemorrhage and exudate may occur. Giant cell eosinophils and arteritis suggest another etiology—allergy. Fibroblastic repair follows. As a rule all stages of activity and healing are seen in the same specimen; occasionally extensive interstitial fibrosis is found alone and is believed to represent the end result of the same process.

In some cases small brownish yellow areas of gelatinous necrosis may be seen with the naked eye particularly in the inner third of the myocardium. The pericardium, endocardium and valves are not involved but mural thrombi are common and may give rise to emboli and infarcts in other organs.

Clinical features. The history is invariably short, rarely longer than a few months. The chief symptoms are increasing dyspnoea and fatigue; some times there is atypical angina pectoris or sub-ternal discomfort (Hansmann and Schenken 1938); occasionally hemiplegia or hæmoptysis signals the onset (Josse and Gallavardin 1901; de la Chapelle and Graef 1931).

The physical signs are usually those of congestive heart failure with a normal or low blood pressure, small pulse pressure, sinus tachycardia, peripheral cyanosis and pallor, cold extremities, general enlargement of the heart (fig. 10.03), gallop rhythm and normal valves. The electrocardiogram often shows left bundle branch block. Angina decubitus may occur.



(a)



(b)

Fig 1002—Focal necrosis in a case of Fiedler's cardiomyopathy

(a) Low power

(b) High power. The cells are macrophages, plasma cells, lymphocytes and eosinophils

(By Dr. J. D. C. H.)



Fig 10 03—Skiagram showing general enlargement of the heart in a case of Fiedler's carditis

although the coronary arteries are healthy. In one such case Bayley (1946) recorded typical anoxic depression of the RS T segment and attributed it to the fact that the lesions were mainly close to the endocardium of both ventricles.

Changes of rhythm are not unusual and include paroxysmal tachycardia, auricular flutter or fibrillation and partial or complete heart block. Ventricular fibrillation and sudden death may occur as in diphtheritic and other forms of myocarditis. Embolic pulmonary infarction may result from dislodgment of right ventricular mural thrombi or from phlebothrombosis in cases of congestive failure. Left ventricular mural thrombi may lead to embolism in the central nervous system, viscera or limbs.

Differential diagnosis. The case usually presents as one of heart failure of uncertain etiology. It is at once distinguished from the hyperkinetic circulatory states (e.g. anaemia, beri beri, arteriovenous aneurysm, Paget's disease of bone, thyrotoxicosis, anoxic pulmonary heart disease, uraemia and certain diseases of the liver) by the obviously low cardiac output, but thyrotoxic heart failure with a low output may cause confusion. Myxoedema should be recognised by the slow heart rate and general features; the basal metabolic rate, electrocardiogram and blood cholesterol will resolve any doubts. Hypertensive pulmonary heart disease may be excluded by the skiagram and electrocardiogram. The heart is usually too large for Pick's disease, but pericardial effusion may be closely simulated; the apex beat is usually felt easily, however, is often forceful and is obviously much displaced to the left; the pulse is not paradoxical and the blood pressure may not be low enough for cardiac tamponade. Gallop rhythm points to a myocardial fault. The electrocardiogram may show left bundle branch block, partial heart block, or RS T and T wave changes more in accordance with myocarditis than with pericarditis. X-ray appearances may be less distinctive, but the right cardiophrenic angle is blunt in heart failure and usually acute in pericardial effusion. Diagnostic paracentesis is advised in doubtful cases.

Rare forms of heart disease which may be confused with isolated myocarditis include congenital hypertrophy, familial cardiomegaly, von Gierke's disease, auricular myxoma, rhabdomyoma, secondary tumours and of course other forms of myocarditis.

Fibrosis of the heart associated with hæmochromatosis or with cirrhosis of the liver from an unbalanced high carbohydrate, low protein diet, should also be borne in mind. The latter occurs particularly in African natives and clinically closely resembles isolated myocarditis. The endocardial fibrosis described by Davies (1948) may have a similar etiology.

Course and prognosis. All proven cases have naturally been fatal, even so there have been no reports of probable or suspected cases which have survived. Death has usually occurred within a few weeks of making the diagnosis or of admitting the patient to hospital.

Treatment. Absolute rest in bed, digitalis, mercurial diuretics, a low

sodium diet and venesection may help to diminish dyspnoea but the general response is poor. Angina decubitus may be relieved by pethidine 50 to 100 mg or phlyseptone, 5 to 10 mg four to eight hourly.

PYOGENIC CARDITIS

In addition to causing toxic myocarditis of the type previously described pyogenic organisms may directly infect the heart. They may produce acute pericarditis with or without sterile or purulent effusion (Chapter XII) or may be responsible for acute bacterial endocarditis (Chapter XI). The staphylococcus and occasionally the pneumococcus may cause miliary abscesses in the heart muscle as part of a general pyæmia. Chemotherapy has greatly improved the prognosis of all forms of pyogenic carditis.

MYOCARDITIS DUE TO DRUGS

Certain therapeutic drugs have earned the reputation of being dangerous to the heart either by causing transient toxic myocarditis or by inducing ventricular fibrillation or asystole. In the first group the best known are digitalis and emetine; in the second chloroform, adrenaline and potassium. Toxic myocarditis due to drug allergy is in a different category and has already been discussed.

DIGITALIS

Digitalis is undoubtedly the best example of a therapeutic drug which may cause dangerous myocardial poisoning.

Pathology. Buchner (1934) first demonstrated that necrotic myocardial lesions could be produced in animals (cats) by means of digitalis. Dearing, Barnes and Essex (1943) also working on cats produced focal necrosis, cellular reaction and fibroblastic repair. Similar necrotic lesions may be provoked by acetylcholine and by continuous direct vagal stimulation (Banting and Hall 1936, 1937) and have been ascribed to coronary constriction. In the belief that the lesions due to digitalis were caused by the activity of acetylcholine, Kyser, Ginsberg and Gilbert (1946) succeeded in preventing them by the simultaneous administration of atropine or a coronary vasodilator such as theophylline. Whether digitalis intoxication in man is characterised by similar patchy myocardial necrosis and whether this is mediated by vagal stimulation remain to be proved but it is a reasonable hypothesis.

Clinical features. Anorexia, nausea or vomiting and diarrhoea usually give sufficient warning of digitalis overdosage but there may be no such indication when carditis from other causes is already present. Disturbances of rhythm are common and include coupling due to premature ectopic beats (fig. 10.04), nodal rhythm, partial or complete heart block (fig. 10.05), multiple ectopic beats, auricular fibrillation, paroxysmal tachycardia (fig. 10.06) and sudden death from ventricular fibrillation.

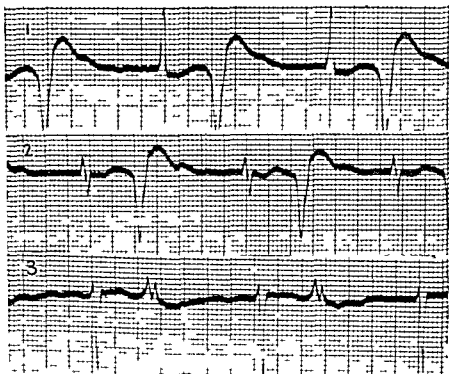


Fig 10 04—Electrocard ogram showing coupling from ventricular ectopic beats due to dig talis



Fig 10 05—Electrocardiogram showing partial heart block due to digitalis

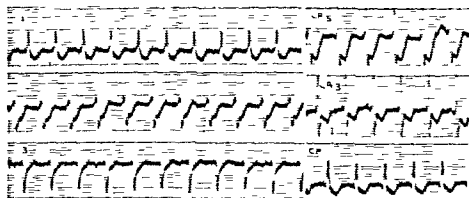


Fig 10 66—Electrocardiogram showing paroxysmal tachycardia due to digitalis.

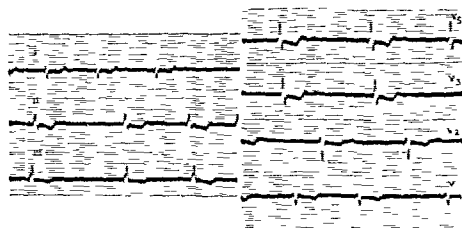


Fig 10 67—Electrocardiogram showing depression of the R-T segment due to digitalis.

The electrocardiogram shows characteristic sagging depression of the RS T segment maximum in leads V₄ 6 when there is normal or increased left ventricular dominance or in leads V₁ 2 when there is right ventricular preponderance. The depression is transmitted chiefly to lead V_L or V_F.

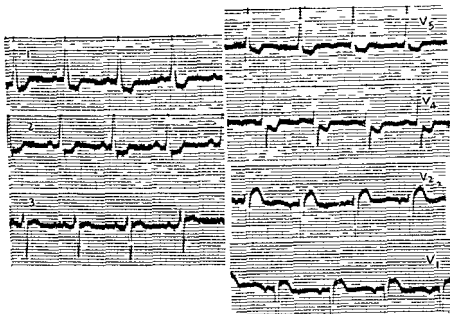


Fig. 10.08—Shortening of the Q-T interval due to digitalis. Q-Tc—0.3 sec

and thence to the appropriate standard lead according to the electrical position of the heart (fig. 10.07). At first the peak of T remains upright but later becomes fused into a more sharply depressed RS-T segment, the Q-T interval being shortened (fig. 10.08). The electrocardiogram offers by far the most reliable evidence of digitalis saturation even when the patient denies having taken the drug.

Treatment. The best remedy, apart from stopping digitalis, is atropine but it is rarely necessary. If the degree of intoxication appears dangerous, however, it may be given in doses of $\frac{1}{2}$ mg. two to four hourly for a day or two.

EMETINE

Emetine is another therapeutic drug with a reputation for causing toxic myocarditis, the chief danger being abnormalities of rhythm, particularly ventricular fibrillation. Emetine was used a great deal amongst British troops in the Mediterranean theatre during the second world war, but ill effects on the heart were very rare if they occurred at all. Patients receiving emetine, however, were always confined to bed throughout the course.

OTHER DRUGS

Potassium when used in large single doses (8 to 16 G) to stop paroxysmal tachycardia or multiple ectopic beats or to differentiate between ischaemic and other causes of T wave inversion is undoubtedly dangerous and may cause sudden death from ventricular asystole, preceded by increasing heart block and bundle branch block (fig 10 09) Spontaneous potassium poisoning may cause sudden death in uraemia

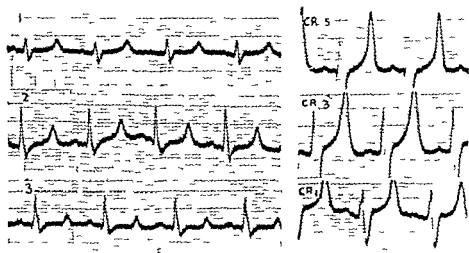


Fig 10 09—Widening of the QRS complex and accentuation of the T wave due to a high blood potassium in a case of uraemia The long Q T is due to hypocalcaemia

Adrenaline in large doses may excite ectopic beats or almost any change of rhythm except heart block Transient hypertension and inversion of the T wave in leads V₄ 6 are common Violent palpitations and substernal discomfort may occur and patients with ischaemic heart disease usually develop a severe attack of angina pectoris Clinical examples may result from errors in the dose of adrenaline administered or from spontaneous hyperadrenalism in cases of pheochromocytoma

Chloroform is an example of a group of drugs mostly anaesthetics which may cause sudden death from ventricular fibrillation especially in the presence of an excess of adrenaline

Nicotine as absorbed by heavy smokers is capable of little more than provoking ectopic beats *Barium chloride* has a similar effect In a minority smoking induces vasoconstriction and may adversely influence hypertension ischaemic heart disease and peripheral vascular disease especially thrombo angitis obliterans It also appears to aggravate thyrotoxicosis

Alcohol is a vasodilator and in moderate amounts may benefit ischaemic heart disease on the other hand it may increase the work of the heart especially if the blood volume is temporarily raised Heavy drinkers may

suffer from an inadequate supply of aneurin and may develop heart failure in consequence (page 514). Finally under the influence of alcohol patients are apt to be careless of medical advice and may exert themselves more than they should. Otherwise there is no evidence that alcohol has any effect upon the heart.

THE HEART IN ACUTE NEPHRITIS

Carditis accompanying acute nephritis (Whitehill *et al* 1939) and toxæmia of pregnancy (Szekely and Sneath 1947) is particularly interesting. The chief clinical features are elevation of the venous pressure, a tendency

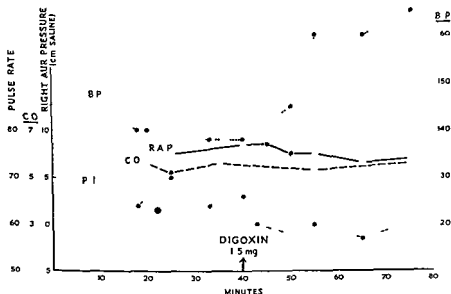


Fig 10 10—Graph illustrating a high right auricular pressure that does not respond to digitalis in a case of acute nephritis. There is a conspicuous rise of blood pressure and slight slowing of the pulse, though cardiac output is unchanged.

to develop acute pulmonary œdema, general enlargement of the heart, and inversion of the T wave in leads facing the surface of the left ventricle (Master, Jaffe and Dack 1937). The degree of hypertension is often insufficient to explain these findings. Nephritic œdema is usually present and the blood volume may be raised.

That there is some form of cardiopathy seems to be proved in certain cases by the behaviour of the cardiac output, which may fail to rise as expected when the venous pressure is high; on the other hand, the lack of response to digitalis (fig 10 10) suggests that the heart is not usually overloaded. Histological examination of the heart muscle in fatal cases of acute nephritis presenting cardiac signs seldom reveals any structural abnormality; sometimes, however, the muscle fibres are dispersed by serous

exudate lymphocytes and endothelial cells — even then there is little if any necrosis (Gore and Saphir 1948)

It is probable therefore that the raised venous pressure is mainly due to an increased blood volume from retention of sodium and water or to some agent causing veno constriction and that as a rule the heart responds normally but that in certain instances cardiac function is impaired owing perhaps to biochemical rather than structural changes in the heart muscle and that acute pulmonary oedema is a manifestation of left ventricular failure even when the blood pressure is but little raised. The subject needs further investigation.

REFERENCES

- Altshuler S S Hoffman K M and Fitzgerald P J (1948) Electrocardiographic changes in diphtheria *Ann Intern Med* 29 294
- Banting F G and Hall G E (1937) Experimental production of myocardial and coronary artery lesions *Tr Ass Amer Phys* 52 204 — — — and Ettinger G H (1936) Experimental production of coronary thrombosis and myocardial failure *Canad med Ass J* 34 9
- Bayles R H (1946) The electrocardiographic effects of injury at the endocardial surface of the left ventricle *Amer Heart J* 31 677
- Bedford D E and Konstam G L S (1946) Heart failure of unknown aetiology in Africans *Brit Heart J* 8 236
- Berry M G Johnson A S and Warshawer S E (1945) Tsutsugamushi fever. Clinical observations in one hundred and ninety five cases *Har Med J* 71
- Buchner F (1934) Herzmuskelnekrosen durch hohe Dosen von Digitalisglykosiden *Arch Exp path Pharmacol* 176 59
- Burch G and Reaser I (1947) A primer of cardiology Philadelphia
- Burkhardt E A Eggleston C and Smith L W (1938) Electrocardiographic changes and peripheral nerve palsies in toxic diphtheria *Amer J med Sc* 195 301
- Chagas C (1909) Nova tripanozomíase humana. Estudos sobre a morfologia e o ciclo evolutivo do *Schizotrypanum cruzi* n. gen. n. sp. agente etiológico de nova entidade morbida do homem *Mem de Inst Osvaldo Cruz Rio de Jan* 1 159
- de la Chapelle C E and Graef I (1931) Acute isolated myocarditis *Arch intern Med* 47 942
- Corbett A J (1943) Scrub typhus *Bull US Army med Dept* 70 34
- Daniels W B *et al* (1943) Meningococcic infection in soldiers *J Amer med Ass* 123 1
- Davies J N P (1948) Endocardial fibrosis in Africans *E African Med J* 25 10
- Dearing W H Barnes A R and Essex H E (1943) Experiments with calculated therapeutic and toxic doses of digitalis effects on myocardial cellular structure *Amer Heart J* 25 648
- Fiedler A (1889) Ueber akute interstitielle Myokarditis. Festschrift zur Feier des 50 Jahr Bestehens des Stadtkrankenhauses zu Dresden Friedrichstadt Dresden part 2 3

French A J (1946) Hypersensitivity in the pathogenesis of the histopathological changes associated with sulfonamide chemotherapy *Amer J Path* 22 679 — Weller C V (1942) Interstitial myocarditis following the clinical experimental use of sulfonamide drugs *Ibid* 18 109

Core I and Saphir O (1947) Myocarditis associated with acute nasopharyngitis and acute tonsillitis *Amer Heart J* 34 831 — — (1947) Myocarditis. A classification of 1402 cases *Ibid* 34 827 — — (1948) Myocarditis associated with acute and subacute glomerulonephritis *Ibid* 36 330

Hansmann G H and Schenken J R (1938) Acute isolated myocarditis *Ibid* 15 749

Howell W I (1945) Absence of electrocardiographic changes in tsutsugamushi fever (scrub typhus) *Arch intern Med* 76 217

Humphreys E M (1948) The cardiac lesions of acute disseminated lupus erythematosus *Ann intern Med* 28 12

Josserand E and Gallavardin L (1901) De l'asthénie progressive des jeunes sujets par myocarditis subaigue primitive *Arch gen de med* 78 513

Kyser F A Ginsberg H and Gilbert N C (1946) The effect of certain drugs upon the cardiotoxic lesions of digitalis in the dog *Amer Heart J* 31 451

Master A M Jaffe H L and Dack S (1937) The heart in acute nephritis *Arch intern Med* 60 1016

Moseley V and Miller H (1945) South American Trypanosomiasis (Chagas Disease) *Ibid* 76 219

Nathanson M H (1928) Electrocardiogram in diphtheria *Ibid* 42 23

Perry C B (1939) Persistent conduction defects following diphtheria *Brit Heart J* 1 111

Porter W B and Bloom N (1935) Heart in typhoid fever: clinical study of 30 patients *Amer Heart J* 10 793

Reinhart W (1946) Isolated diffuse interstitial eosinophilic myocarditis *Cardiologia* 11 219

Scott R W and Saphir O (1929) Acute isolated myocarditis *Amer Heart J* 5 129

Szekely P and Sneath L (1947) The heart in toxæmia of pregnancy *Brit Heart J* 9 128

White P *et al* (1937) Heart 15-20 years after diphtheria *Amer Heart J* 13 534

Whitehill M R Longcope W T and Williams R (1939) The occurrence and significance of myocardial failure in acute hæmorrhagic nephritis *Bull Johns Hopk Hosp* 64 83

Williams S W Sinclair A J M and Jackson A V (1944) Mite borne (scrub) typhus in Papua and the Mandated Territory of New Guinea. Report of 66 cases *Med J Australia* 2 525

Wood P H (1941) Differential diagnosis of Da Costa's syndrome *Proc Roy Soc Med* 34 543

BACTERIAL ENDOCARDITIS

BACTERIAL or infective endocarditis means bacterial infection of any of the heart valves or of certain congenital anomalies of the heart or great vessels (bacterial endarteritis). It occurs in two main forms acute (malignant) due to infection with any of the pyogenic bacteria and subacute due mainly to the *Streptococcus viridans* but many other organisms have been isolated from both types. This broad classification is necessarily artificial the course of the disease depending on the virulence of the organism and the resistance of the host. There is no clear division between the two types and they are better considered as one disease.

There is usually some underlying fault congenital or acquired. The most susceptible congenital anomalies are pulmonary stenosis bicuspid aortic valve ventricular septal defect and patent ductus arteriosus atrial septal defect is remarkably immune. Any acquired valve lesion may become infected including syphilitic aortic incompetence (Martin and Adams 1938) and calcific aortic stenosis (Brink and Smith 1937) but old rheumatic valvulitis is commonly to blame. In quite a number active rheumatic infection is still present when bacterial endocarditis is superimposed.

PATHOLOGY

The lesion is superficial and is not a valvulitis in the sense that rheumatic endocarditis is bacteria invade the surface of a damaged or congenitally deformed valve and are encouraged by the formation of small superficial thrombi which provide an excellent culture medium. Both in the natural disease and experimentally in dogs there appears to be a paucity of granulation tissue and of cellular reaction the microbes are not destroyed and healing does not take place. Elsewhere in the body similar foci of bacteria are rapidly walled off by granulation tissue and the lesion is invaded by leucocytes the microbes are destroyed and the inflammation soon subsides (Friedman Katz Howell *et al* 1938).

The macroscopic appearances vary according to the infecting organism tending to be finely granular with streptococcus viridans ulcerative and hæmorrhagic with the hæmolytic streptococcus and pneumococcus proliferative with the gonococcus. When associated with congenital defects the site of the vegetations depends upon the direction of blood flow through the defect thus in the maladie de Roger vegetations are found on the right side of the patent interventricular septum and on the wall of the right ventricle opposite the defect with patent ductus arteriosus they are found

at the pulmonary artery end. Ulceration may lead to perforation of a valve cusp or sinus of Valsalva. In old rheumatic cases vegetations may spread on to the endocardium of the left auricle (Thayer 1926).

The myocardium may show scattered focal lesions similar to those seen in isolated or toxic myocarditis or small collections of lymphocytes or lymphocytes and polymorphs known as Bracht-Wachter bodies (Bracht and Wachter 1909). The latter are believed to be embolic in origin and represent a local inflammatory reaction to bacterial nests (Perry 1936). They are the non-suppurative counterpart of the miliary abscesses seen in staphylococcal cases.

OCCURRENCE

Bacterial endocarditis accounts for about 2 per cent of all cases of organic heart disease (White 1937). It may occur at any age but is most common in young adults. Males are affected rather more frequently than females. Auricular fibrillation occurs in only 2.5 per cent of cases (McDonald 1946) presumably because it is not a feature of the congenital lesions mentioned. It is uncommon in rheumatic aortic valve disease and occurs late in the life history of patients with mitral stenosis. There is no evidence that the two conditions are mutually antagonistic.

CLINICAL FEATURES

Patients may present themselves with cardiac symptoms: pyrexia of unknown origin, anaemia, a cerebral vascular lesion, subacute rheumatism, nephritis, broncho-pneumonia or with other patterns which depend upon the nature of the invading organism, the underlying cardiac lesion and the caprice of the disease process. At the onset symptoms are often ascribed to influenza but fail to clear up. The diagnosis rests upon the combination of a variety of signs which will be considered individually.

Cardiac abnormalities. Evidence may be present of one or other of the various underlying valve lesions or congenital defects already mentioned or there may be just an impressive systolic murmur at the mitral area but if there are no abnormal auscultatory signs of heart disease the diagnosis is rarely tenable. The development of a new valve lesion or of the whining diastolic murmur and thrill of a perforated aortic cusp may be highly suggestive.

Toxic myocarditis is not uncommon and may cause heart failure and death whether the infection yields to treatment or not. Its importance has been more widely recognised since the introduction of penicillin.

Pyrexia. Acute cases are always febrile; subacute cases are always febrile at some stage in the disease but bouts of fever may alternate with afebrile periods. The fever is irregular in type, usually low grade or moderate in degree and may continue for weeks, months or years.

Anaemia. Anaemia nearly always develops early and is already present in

about three quarters of the patients when first seen. It is indeterminate in type being normocytic and isochromic even when associated with hæmolytic infections. The red cells may be reduced to about three million and the hemoglobin to about 60 per cent giving a normal colour index. Stained films and bone marrow samples reveal no specific features. If microcytic hypochromic anæmia is found the diagnosis should be doubted for iron deficiency anæmia itself may cause many of the signs and symptoms of bacterial endocarditis e.g. functional systolic murmurs at the base or the apex of the heart splenomegaly petechiæ red cells in the urine and even low grade pyrexia.

The white count is variable. It may be normal on the other hand there may be moderate leucocytosis or leucopenia. Leucocytosis is usually associated with acute septicæmic cases normal or leucopenic counts with subacute infections.

Splenomegaly. The spleen is usually palpable. It may be soft as in typhoid when due to septicæmia it may enlarge rather suddenly as a result of splenic infarction when it is tender it may be firm in subacute cases or it may be so large as to cross the mid line in chronic cases.

Petechiæ. Petechiæ are common and sometimes appear in successive crops. They may be seen under the nails in the ocular fundi in the conjunctivæ or anywhere in the skin or mucous membranes. Under the nails they resemble small splinters (Horder 1926) in the fundi they may have white centres of exudate in the skin they must be distinguished from minute telangiectases—Campbell de Morgan's spots. Petechiæ in successive crops or otherwise are in no way diagnostic of bacterial endocarditis. They are due to capillary hæmorrhage and may occur in any condition in which the capillaries are suitably damaged including most forms of septicæmia acute rheumatic fever (especially when associated with acute glomerulonephritis) and severe anæmia. In bacterial endocarditis the capillary lesion may be due to toxins to allergy or to anæmia.

Increased capillary fragility may be demonstrated by the capillary resistance test.

A cuff is placed on the upper arm inflated to a pressure of 50 mm. of mercury and maintained for five minutes alternatively a pressure of 80 mm. of mercury may be maintained for three minutes. The arm below the cuff is then inspected. Most normal subjects are unaffected but some develop a few tiny petechiæ in the antecubital fossa. The result of the test may be expressed as slightly moderately considerably or grossly positive or as negative the four positive grades representing transition from a few tiny hæmorrhages to gross purpura.

It may be positive or negative in bacterial endocarditis when spontaneous petechiæ are present. When positive it is well to make sure that vitamin deficiency is not responsible or to cover this possibility by giving adequate doses of ascorbic acid rutin and crude vitamin P.

Small hæmorrhagic pustules in the skin may occur in the acute pyogenic forms of bacterial endocarditis and are embolic in origin.

Clubbing of the fingers (and toes) Clubbing occurs in about half of the subacute cases but as it takes at least 3 to 6 weeks to develop it is rare in malignant endocarditis. Early clubbing may be recognised by noting congestion and thickening of the nail fold and loss of the normal angulation between the nail fold and the base of the nail. Slight clubbing should be interpreted with caution however for it may occur in many conditions including active rheumatic carditis. Conspicuous clubbing on the other hand provides excellent supportive evidence of bacterial endocarditis if cyanotic congenital heart disease pulmonary abscess bronchogenic carcinoma and a congenital origin can be excluded. The mechanism of clubbing is not yet fully understood.

Nodes Osler's nodes are small transient erythematous lesions about the size of a pea lasting a few days and vivid pink in colour when fresh bluish when fading often with a darker centre they are raised palpable and tender and may be found particularly on the pads of the fingers and toes on the sides of the fingers or on the thenar or hypothenar eminences (Osler 1909). They are due to infected cutaneous emboli and the responsible organism may sometimes be cultured from them.

More important perhaps because more common are larger deeper nodes which vary from the size of a pea to that of a grapefruit. They are red painful hot and tender may occur anywhere in the limbs and may be mistaken for osteomyelitis or periostitis. When a lesion involves the finger it closely resembles an ordinary infected pulp it is non suppurative however and disappears in about a week if left alone. Cultures from the inflamed tissue may yield *Streptococcus viridans*. Red tender macules are equally characteristic and even more common and may also yield positive cultures from biopsies.

Emboli In addition to the minute emboli which cause white centred petechiae and the nodes just mentioned larger emboli may block any artery—cerebral visceral or peripheral. They are more common in the radial ulnar posterior tibial and dorsal artery of the foot than in the axillary or femoral artery because their size is limited. For this reason peripheral emboli are often symptomless and are only discovered by those who look for them. In cases of suspected bacterial endocarditis the peripheral vessels should always be palpated and their patency noted for future reference.

Mycotic aneurysm Ulceration or degeneration of the wall of an artery due to local inflammation from an infected embolus lodging within the vessel or in its vasa vasorum may result in the formation of a small aneurysm. In a peripheral vessel this is controllable but when it occurs in a visceral artery fatal hæmorrhage from spontaneous rupture may ensue.

Pulmonary emboli When bacterial endocarditis involves the pulmonary or tricuspid valve or when it is associated with a left to right cardiac shunt as in patent interventricular septum emboli may be flung into the pulmonary circulation. Numerous small pulmonary infarcts result and may

give rise to a clinical picture resembling recurrent or subacute hæmorrhagic bronchopneumonia

Renal lesions The various renal lesions that may occur in bacterial endocarditis represent almost every aspect of the disease

(1) An embolus lodging in a small renal artery leads to simple infarction of the kidney with hæmaturia or without signs or symptoms

(2) Minute bacterial emboli may cause embolic nephritis which in greater or less degree is found in the majority of cases. Only some of the glomeruli are involved, rarely more than 60 per cent and most of these have some of their capillary loops intact so that the tuft is not entirely avascular and the health of the tubules is not seriously threatened. Affected capillaries are converted into a hyaline mass and red cells may be found in the capsular space and in the urine. Embolic nephritis does not cause renal failure because a sufficient number of glomeruli are always spared (Baehr 1921)

(3) In acute pyogenic forms of bacterial endocarditis particularly when pneumococcal or staphylococcal in origin multiple abscesses may be found in the substance of the kidney

(4) Petechiæ due to simple capillary hæmorrhage may occur on the surface of the kidney in the absence of embolic nephritis. They are then similar to those found in the pericardium, pleura and skin

(5) Ordinary acute diffuse glomerulo nephritis may occur as with other streptococcal infections and may progress to renal failure but not more than 5 to 10 per cent of all cases take this course

(6) Simple congestion of the kidney may result from heart failure and give rise to albuminuria and to a few red cells in the urine

It will be appreciated that these six types of renal lesion represent thrombotic emboli, benign bacterial emboli, septic emboli, simple hæmorrhage, toxæmia or allergy and heart failure respectively and that nearly all the features of bacterial endocarditis may be understood in terms of these six factors

Changes in the ocular fundus Simple petechiæ like those in the skin are fairly common. Occasionally, they have white centres and may be embolic in origin. It should be understood that these white centres represent exudate and that identical lesions may be seen in other conditions particularly leucæmia, pernicious anæmia and malignant hypertension. The exudate may be surrounded by hæmorrhage or may be to one side of it. Embolism of the central artery of the retina or of one of its main branches may cause complete or partial loss of vision but is fortunately rare. Finally papilloedema or papillitis with or without widespread hæmorrhages and exudates is not uncommon when there is diffuse glomerulo nephritis the appearances resembling those of malignant hypertension

DIAGNOSIS

It is emphasised that pyrexia anæmia splenomegaly petechiæ and diffuse glomerulo nephritis may occur wherever the site of the cardiac lesion that systemic emboli mycotic aneurysms nodes and embolic nephritis signify left sided lesions e.g. aortic or mitral valve disease that multiple hæmorrhagic infarcts in the lungs are the prerogative of right sided valve lesions and of left to right congenital shunts such as patent ductus and maladie de Roger (Barker, 1949)

Clinically bacterial endocarditis should be considered in all cases of unexplained fever with suspicious auscultatory signs in the heart. If an indeterminate anæmia is also present a determined search should be made for other evidence. If splenomegaly petechiæ and red cells in the urine are added the diagnosis becomes probable but is still uncertain. On the other hand clubbing of the fingers nodes peripheral emboli mycotic aneurysm nephritis and characteristic fundal changes may each one of them be diagnostic of bacterial endocarditis when associated with fever and an appropriate cardiac lesion.

The diagnosis is confirmed by a positive blood or bone marrow culture. Six tubes are usually set up from each sample and 4 to 6 samples should be obtained at different times preferably when the temperature is high before a negative result is accepted. It should be pointed out however that blood cultures from patients with pyorrhœa or with dental abscess may grow *Streptococcus viridans* when the specimen is obtained after chewing so that the diagnosis of bacterial endocarditis should never rest on a positive blood culture alone.

NATURAL COURSE

Untreated patients with acute infection die in a matter of days or weeks usually from septicæmia or from the effects of embolism. Those with subacute infection usually live for months and occasionally for years. Bouts of fever with exacerbation of signs and symptoms alternating with afebrile quiescent phases described by Libman as bacteria free periods. Death may result from heart failure cerebral or other visceral embolism hæmorrhage uræmia or other causes. According to Libman and Friedberg (1941) about 3 per cent of all patients recover spontaneously but Lichtman (1943) found that only 1 per cent of 2 596 cases collected from the literature so recovered.

PROGNOSIS

Penicillin and streptomycin have radically altered the course of bacterial endocarditis for the infection can now be controlled in 90 per cent of cases. However about 35 per cent still die during or shortly after treatment mostly from heart failure. This high mortality may be due to the frequency of serious toxic myocarditis and to the relatively rapid increase in severity

of valve lesions. Uræmia accounted for only 6 per cent of the 131 deaths in the combined hospitals series reported by Christie (1948) emboli for 11 per cent and hæmorrhage for 8 per cent. The most important factors influencing the mortality rate proved to be the presence and degree of heart failure, the duration of the infection, and the nutritional state of the patient.

Relapses are common in inadequately treated cases, but should not exceed 10 per cent in patients who have received at least 0.5 mega unit of penicillin daily for a minimum period of 28 days. Nearly all those who relapse do so within one month of ceasing treatment. The frequency of recurrence (as distinct from relapse) is not yet known.

TREATMENT

Prophylactic. Surgical repair of patent ductus arteriosus not only cures the defect, but protects the patient from infective endarteritis. Repair of coarctation of the aorta may be less successful in the second respect, because infection may yet complicate an associated bicuspid aortic valve.

Dental hygiene is particularly important in all patients who have congenital heart disease or chronic valve disease. Tooth extractions, tonsillectomy, and other I.N.T. operations should be covered by 100,000 units of penicillin six hourly for 48 hours.

Acute pyogenic infections should be treated with penicillin rather than sulphonamides in patients who are susceptible to bacterial endocarditis.

Chemotherapy. Sulphonamides have proved disappointing, and although they may temporarily sterilise the blood stream and lower the temperature, they rarely cure the disease. Chemotherapy is at a disadvantage because bacterial endocarditis is practically avascular. As previously stated, bacteria are buried in a mass of thrombotic vegetations which serve as an excellent culture medium. It is reasonable to suppose that if the formation of such thrombi could be prevented, the culture medium would become exhausted and the bacteria would starve or become exposed to chemotherapy. To effect this, heparin has been administered intravenously, either by the drip method or by four hourly injection, over a period of weeks, the dosage being controlled by the clotting time of the blood which is kept at 20 to 60 minutes.

I tried this treatment for the first time in 1937 in an early case of bacterial endocarditis due to *Streptococcus viridans* on the mitral valve.

The patient was a youngish woman in a good state of nutrition, who presented with organic mitral incompetence, continued pyrexia of moderate degree, splenomegaly, and repeated positive blood cultures. As she had been told the disease was fatal, she willingly submitted to the experiment. Heparin was given by intravenous injections at four hourly intervals over a period of three weeks, and the clotting time was kept between half an hour and four hours. From time to time she bled rather profusely from her teeth, but there were no other ill effects until the end. In addition to the heparin, a full course of sulphanilamide was adminis-

tered from the tenth day onwards. Two hours after the last dose of heparin when her clotting time was still in the region of one hour she died suddenly of subarachnoid hemorrhage due possibly to a ruptured mycotic aneurysm. Owing to the prolonged clotting time she had no chance against this catastrophe. At autopsy (when the blood was still unclotted) old vegetations were seen on the mitral valve but there were no recent thrombi the surface of the granulations being smooth and white. Macroscopically the object of the treatment appeared to have been achieved. Microscopically however numerous bacteria could be seen in the old thrombi cultures from which were positive.

It was concluded from this experiment which offered a peculiar opportunity for autopsy inspection at the critical moment that three weeks heparin treatment combined with sulphanilamide was ineffective. Subsequent work has confirmed this observation (Leach *et al.* 1941).

Another method which it was hoped would yield more fruitful results was to combine chemotherapy with hyperthermia for sulphonamides are more potent at higher temperatures *in vitro* (White and Parker 1938) but this proved little better than heparin. Of a series of 489 cases treated with sulphonamides alone collected by Lichtman (1943) 4 per cent recovered of 109 cases treated with sulphonamides combined with heparin or hyperthermia 6.4 per cent recovered.

The situation has greatly improved since the introduction of penicillin. Patients should be treated early as soon as the diagnosis is clinically probable without waiting for positive results of blood cultures. Every effort should be made to counter malnutrition and a blood transfusion should be given if there is serious anaemia.

The standard dose of penicillin is 0.5 mega unit daily given in divided doses of 60 000 units three hourly, 80 000 units four hourly or 120 000 units six hourly and continued for twenty eight days. Nothing less than this will suffice and larger doses prolonged for six to eight weeks are often necessary. One of my patients was not controlled until she received a million units three hourly and a total of 250 million units. If the resistance of the organism is known so much the better but even then the optimum dose cannot be calculated exactly because it depends partly on the physical properties of the lesion. Swift and maintained clinical response is the only reliable criterion by which to judge the correct dose. If however the resistance of the organism is known to be more than eight times that of the standard test strain of Oxford staphylococcus the dose of penicillin should certainly be increased proportionately (Christie 1948). If the coefficient of resistance is 10 not less than 100 000 units three hourly will suffice if 20 then at least 200 000 units three hourly will be necessary—and so on (Baehr and Gerber 1947). Peak (15 to 30 minutes after intramuscular injection) or constant (with the intravenous drip method) blood serum levels of penicillin expressed in units per ml may also be measured and checked against tables giving the expected level for the dose employed. Peak levels should range from 2 to 25 units per ml with doses of 60 000 to

500 000 units intramuscularly constant levels between 1 and 10 units per ml with daily doses of 500 000 to 4 million units

To avoid the discomfort of frequent needling there is an increasing tendency to give massive doses of penicillin (0.25 to 0.5 mega unit) three or four times daily. As these massive doses have a penetrating power denied to more modest quantities there is something to be said for this method but they should not be given too infrequently.

Another way of reducing the frequency of injections while maintaining a sufficiently high blood level over the 24 hours is to give a suspension of procaine penicillin in oil (Herrell, Nichols and Heilman, 1947) ampoules are available containing 300 000 units per ml. injections are painless and may be given once daily. Particular care should be exercised in using oily solutions on patients with right to left intracardiac shunts owing to the danger of paradoxical cerebral embolism.

Finally the blood level of penicillin may be increased up to fourfold by the oral administration of certain substances such as benzoic acid (Bronfenbrenner and Favour, 1945) sodium benzoate or caronamide (4^1 /carboxy/phenylmethane sulphonamide) which interfere with penicillin excretion by the renal tubules. The dose of each of these substances is 2-3 G four-hourly (Boger *et al.* 1948). Caronamide may be combined with sodium benzoate with some advantage and is a valuable adjunct to treatment in highly resistant cases.

Treatment of relapses or resistant cases. If the previous course of treatment was inadequate in dosage or duration the standard 28 day course should be instituted but if a relapse follows adequate treatment every effort should be made to culture the organism and to determine its sensitivity to penicillin. If its resistance is not greater than eight times the standard the dose of penicillin should be doubled and treatment should be continued for six weeks. If the coefficient of resistance is greater than 8 the dose of penicillin should be increased proportionately. If the organism is highly resistant or if it has not been isolated and the infection remains uncontrolled streptomycin may be tried. The dose is 1 to 3 G daily for four to six weeks. Caronamide does not influence the blood level of streptomycin for the latter is excreted by the glomeruli.

Toxic reactions of penicillin. Apart from local pain from subcutaneous or superficial intramuscular injections and phlebotrombosis from intravenous injections the only toxic manifestations which can be attributed to penicillin are fever and urticaria. Fever was common when crude penicillin was used but is rarely seen nowadays. Urticaria develops in about 5 per cent of cases and may be extreme soft tissue œdema and hydrarthrosis are occasionally associated. This allergic reaction is alleviated by adrenaline and by the antihistamine group of drugs. Penicillin may be continued in mild cases but may have to be stopped if the reaction is severe.

The chief toxic effect of streptomycin is on the vestibular

as of

the sense of balance may be permanent if heavy doses are continued after giddiness has developed. A conservative dose of 1 G daily however may be continued in the presence of minor vestibular symptoms if the latter are controlled by means of antihistamine drugs.

Less toxic antibiotics such as chloromycetin are likely to replace streptomycin in the near future.

Other considerations An infected ductus should be controlled by penicillin then ligated as soon as the patient is fit enough.

Heart failure should be treated in the customary fashion but the prognosis is grave in these cases.

Diffuse glomerulo nephritis may be mistaken for a relapse of bacterial endocarditis. If the renal function is impaired the blood level of penicillin may rise considerably and thus aid the primary treatment unfortunately however the nephritis usually proves fatal.

REFERENCES

- Bachr G (1921) The significance of the embolic glomerular lesions of subacute streptococcus endocarditis. *Arch intern Med* 27 262 — and Gerber I E (1947) Penicillin treatment of subacute bacterial endocarditis. *Advances intern Med* 2 308.
- Barker I S (1949) A clinical study of subacute bacterial infection confined to the right side of the heart or the pulmonary artery. *Amer Heart J* 37 1054.
- Boger W P, Miller A K, Tillson E K and Shaner G A (1948) Caronamide plasma concentrations, urinary recoveries and dosage. *J lab and clin Med* 33 297.
- Bracht E and Wachter (1909) Beitrag zur aetologie und pathologischen anatomie der myokarditis rheumatica. *Deut Arch f klin Med* 96 493.
- Brink J R and Smith M L (1937) Subacute bacterial endocarditis: clinico-pathological study of 37 cases. *Amer Heart J* 14 362.
- Bronfenbrenner J and Favour C B (1945) Increasing and prolonging blood penicillin concentrations following intramuscular administration. *Science* 101 673.
- Christie R V (1948) Penicillin in subacute bacterial endocarditis. *Brit med J* 1 1.
- Friedman M, Katz L N, Howell K M, Lindner E and Mendlowitz M (1938) Experimental endocarditis due to streptococcus viridans. *Arch intern Med* 61 95.
- Herrell W E, Nichols D R and Heilman F R (1947) Procaine penicillin C (duracillin): a new salt of penicillin which prolongs the action of penicillin. *Proc Mayo Clinic* 22 567.
- Horder T (1906) Lumsden lectures on endocarditis. *Lancet* i 695, 745, 850.
- Leach C E (1941) Chemo therapy and heparin in subacute bacterial endocarditis: further experiences. *J Amer med Ass* 117 1345.
- Libman E and Friedberg C K (1941) Subacute bacterial endocarditis. New York.
- Lichtman S S (1943) Treatment of subacute bacterial endocarditis: current results. *Ann intern Med* 19 787.
- McDonald R K (1946) The coincidence of auricular fibrillation and bacterial endocarditis. *Amer Heart J* 31 308.
- Martin H E and Adams W L Jr (1938) Bacterial endocarditis superimposed on syphilitic aortitis and valvulitis. *Ibid* 16 714.

Osler W (1909) Chronic infectious endocarditis *Quart J Med* 2 219

Perry C B (1936) Bacterial endocarditis Bristol

Phayer W S (1926) Studies on bacterial (infectious) endocarditis *Johns
Hopk Hosp Rep* 12 1

White H J and Parker J M (1938) Bacterial effect of sulphanilamide upon
beta hæmolytic streptococci in vitro *J Bact* 36 481

White P D (1937) Heart disease New York

CHAPTER VII

PERICARDITIS

THE features of pericarditis depend upon its etiology upon the presence or absence of effusion upon the nature and hydrostatic pressure of such effusion and upon the development or otherwise of constriction in chronic or adhesive cases

ETIOLOGY

Pericarditis may be rheumatic tuberculous pyogenic traumatic uræmic or secondary to myocardial infarction malignant growths may invade the pericardium hæmopericardium may result from rupture of a syphilitic or dissecting aneurysm from perforation of a myocardial infarct or ventricular aneurysm or from stab or gun shot wounds of the heart hydropericardium may complicate congestive heart failure or myxœdema some times the etiology is obscure All these types have their own special characteristics which will be described subsequently but they have also certain features in common

DRY (FIBRINOUS) PERICARDITIS

All varieties of pericardial inflammation may present in this form The diagnosis rests on three cardinal signs pain pericardial friction and a specific electrocardiographic pattern Disturbances of temperature pulse rate sedimentation rate etc of course may occur but are of little help in diagnosis

Pain (apps (1932) found that the pericardium was insensitive to stimuli calculated to produce pain except in that part of it roughly its lower third which is supplied by the phrenic nerve It follows that pericarditis should be painless unless the pain has phrenic distribution or unless it is pleural in type from secondary involvement of that structure In fact this is so in the majority of cases there is no pain in some pain is referred to the neck or shoulder tip in others it is precordial and pleural in type catching the breath on inspiration or on coughing

Pericardial friction Friction sounds may be heard anywhere over the heart according to the site and nature of the pathological process but are most common at the left sternal border in the fourth intercostal space over the area of maximum cardiac dullness where the pericardium lies in contact with the chest wall They are superficial rough or smooth loud or soft their timing is peculiar being out of step with the heart sounds Some times they are confused with the to and fro murmur of aortic valve disease

or with artificial stethoscopic sounds sometimes they escape detection. Pleuro pericardial friction can usually be distinguished by its relationship to respiration.

Electrocardiographic changes A diagnostic electrocardiographic T pattern, first described by Porte and Pardee (1929) may be found in the majority of cases of genuine pericarditis whatever the etiology, and whether

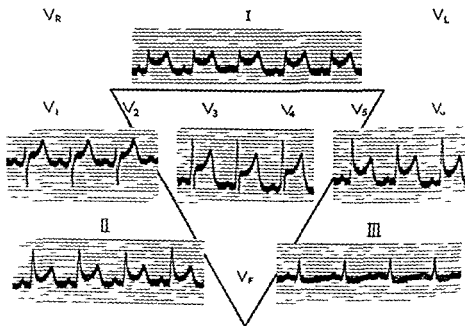
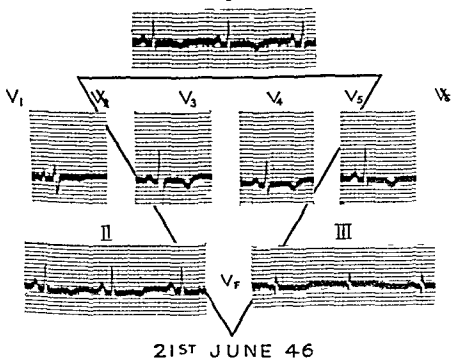


Fig. 12.01—Electrocardiogram showing the early phase of the pericardial T₂ pattern. This graph is atypical in that the R-T segment is not elevated in lead 3.

or not there is effusion (Wood, 1937). It develops in two stages, early and late, the changes usually appearing in all leads and therefore especially in lead 2. In the early phase (fig. 12.01) the RS-T segment is elevated but retains its natural concavity. Within a few days it regains the iso-potential level or becomes depressed, and the T wave becomes flattened, diphasic or inverted (fig. 12.02). QRS remaining unchanged throughout or losing voltage. When the inflammation subsides the graph returns to normal except when a tuberculous or pyogenic pericarditis merges into the chronic constrictive form, when flat or inverted T waves and low voltage QRS complexes become permanent. The T pattern may only be appreciated in serial electrocardiograms, as changes may be confined to leads 1 and 2 in one record and to leads 2 and 3 in another. Similar appearances are seen in all chest leads and may be found when limb leads are normal (fig. 12.03).

Both early and late stages appear to be due to alterations in the biophysical properties of the sub-epicardial myocardium, whether or not there



I

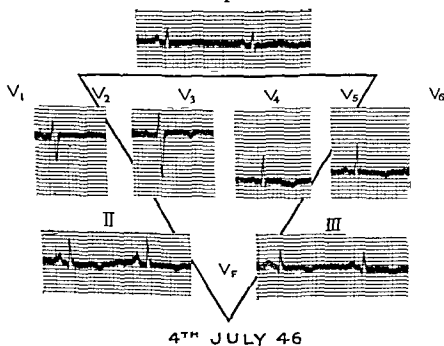


Fig 02—Electrocardiogram showing the presence of the pathological T₂ pattern characteristic of pyogenic pericarditis secondary to bronchopneumonia

are recognisable structural changes (Kisch *et al* 1940). The early pattern of generalised pericarditis may be distinguished from that of myocardial infarction by the absence of a conspicuous Q wave by the preservation of the upward concavity of the RS T segment and by the occurrence of maximum RS T deviation in lead 2 (cf T_1 and T_3 types in myocardial infarction). When pericarditis is localised however changes may be maximum in leads 1 or 3 (Burchell Barnes and Mann 1939). The later stage

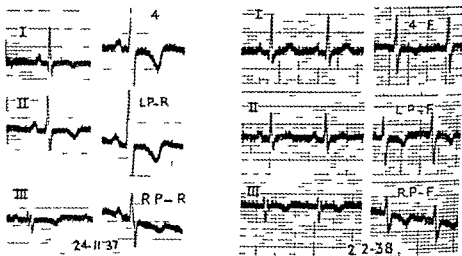


Fig 1203—Electrocardiogram showing late changes due to pericarditis in the second record (2nd February 1948) they are limited to the chest leads

may be confused with active carditis, myxoedema, carbon monoxide poisoning, severe anaemia, concordant left ventricular preponderance, combined anterior and posterior myocardial infarction, and long standing congestive heart failure from any cause. On the other hand, the characteristic initial phase, the changing picture in serial graphs, and the clinical features of the case usually make the diagnosis easy.

PERICARDIAL EFFUSION

Fluid in the pericardial sac may be a simple transudate (hydropericardium), a straw coloured sterile exudate, a purulent exudate, or blood (haemopericardium). It may disturb the patient in one or more of four ways: (1) stretching of the pericardium may induce praecordial discomfort; (2) large effusions exerting pressure on surrounding structures, especially on the bronchi and lungs, may produce reflex cough and dyspnoea; (3) if the fluid is purulent, there may be constitutional effects similar to empyema; (4) as the pressure rises in the pericardial sac, cardiac filling is hampered, the pressure rises in the systemic veins, the ventricular stroke output diminishes, and the blood pressure tends to fall. The raised venous

pressure is partly beneficial for it aids cardiac filling the diminished stroke volume is countered by tachycardia vasoconstriction serves to maintain the blood pressure (Stewart Crine and Deitrich 1938) When these compensatory adjustments fail to meet the circulatory demands the situation becomes critical (cardiac tamponade)

Effusions in excess of 250 ml may be detected by percussion Dullness may be elicited in the second left space when the patient lies flat to the left of the apex beat when the latter can be located in the xiphisternal angle and to the right of the sternum in the 4th and 5th intercostal spaces (Rotch's sign 1878)

Auscultation reveals pericardial friction in the majority of instances even with gross effusions and the heart sounds are faint Signs of pulmonary collapse are common at the left base (Ewart's sign 1896) and consist of dullness to percussion increased tactile fremitus bronchial breathing and whispering pectoriloquy with no adventitious sounds

Fluoroscopy shows the natural contours of the heart to be obliterated by a large globular or pear shaped shadow which may change its shape with alteration of posture The right border usually meets the diaphragm at an acute angle (fig 12 04) in contrast to the blunt right cardiophrenic angle of tricuspid valve disease (fig 12 06) Pulsation is diminished and may be absent The size of the shadow may change rapidly and grossly from week to week (figs 12 04 and 12 05) In the first oblique position the concave line of the inferior vena cava at the posterior inferior angle of the heart shadow is replaced by a convex backward bulge

Cardiac tamponade The pressure within the pericardial sac depends upon the quantity of fluid the rate at which it accumulates and the elasticity of the parietal pericardium The presence and degree of cardiac compression or tamponade may be assessed clinically by estimating the blood pressure the venous pressure the pulse rate and the amount of vasoconstriction

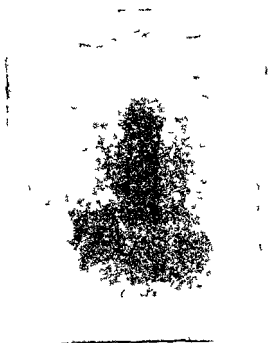


Fig 12 04—Skiagram of a case of pericardial effusion showing an acute right cardiophrenic angle



Fig 1-05- Skuagram showing rapid diminution in size of cardiac silhouette as effusion is absorbed Taken six weeks after Fig 12-04



F g 2 06—bk ag am showing the blunt ght c d o ph en c angle in tricusp d ncom
p ten e

Paracentesis is advised if the blood pressure falls below 90 mm. of mercury or if there is a high venous pressure together with marked tachycardia and conspicuous vasoconstriction. The cervical veins pulsate less than in congestive heart failure and considerably less than in tricuspid incompetence. Vasoconstriction results in pallor, cyanosis and coldness of the skin especially of the extremities.

There is reason to believe that cardiac tamponade seriously interferes with the coronary blood flow, not only because the cardiac output is reduced and the blood pressure low, but because the pressure gradient between the aorta and coronary circulation is significantly reduced. The myocardium appears to suffer accordingly and true heart failure may result. This may explain those cases that fail to recover after decompression and suggests that tamponade should be regarded as a medical emergency.

Differential diagnosis. It may be by no means easy to distinguish pericardial effusion from acute dilatation of the heart clinically or radiologically. The electrocardiogram may help, Q-Tc being relatively short in pericardial effusion and long when the heart is dilated. When the diagnosis is in doubt, however, it is best to explore with a needle for the treatment of the two conditions is radically different.

Treatment. The object of treatment is to prevent death from cardiac tamponade and is achieved by avoiding therapeutic agents which may lower the venous pressure, such as mersalyl, a low sodium diet and venesection, and by decompression if necessary. Paracentesis may be carried out to the left of the apex beat or at any point where there is reason to believe there is plenty of fluid. If the needle touches the heart, forcible pulsation can be felt and it should be withdrawn a little or inserted elsewhere, with due care the risk of causing hæmopericardium from puncturing a coronary vessel is small. Fluid may also be removed if purulent or if untoward symptoms are caused by pressure on surrounding structures. Hæmopericardium, which is responsible for many cases of tamponade, requires surgical evacuation and repair of the underlying injury.

CHRONIC CONSTRICTIVE PERICARDITIS

Although Richard Lower described the paradoxical pulse and calcified pericardium as early as 1669, he was not in a position to grasp their full significance and it was Chevers who really drew attention to the disease, giving an excellent account of it with considerable understanding of the circulatory dynamics involved in 1842. The term 'Pick's disease' is unfortunate for Pick (1896) merely emphasised the accompanying pseudo-cirrhosis of the liver and because priority undoubtedly goes to Chevers. The issue is best avoided by adhering to the descriptive title—chronic constrictive pericarditis.

Morbid anatomy. The condition may be regarded as a complication of the healing process following tuberculous, pyogenic and perhaps certain other forms of pericarditis, for the fibrous tissue which may be

hid down so extensively in the active phase of these diseases contracts on maturation and limits diastolic expansion of the heart. Calcium is often deposited in large quantities and the whole heart may become encased in stone.

Etiology. Tuberculosis accounts for at least three quarters of the cases and may be still active when constriction first develops. The pyogenic bacteria appear to be responsible for a few and the cause is uncertain or unknown in the remainder. None are rheumatic (White 1935).

Clinical features. The clinical picture is one of more or less generalised cardiac compression, the circulatory dynamics being somewhat similar to those of high pressure pericardial effusion. The unyielding pericardium however sets a rigid limit to maximum cardiac filling, no matter how high the venous pressure and the cardiac output can only be increased by acceleration of the pulse (Stewart and Heuer 1939). Patients usually present themselves with dropsy and ascites and may feel relatively well. They may complain of breathlessness on exertion but are usually comfortable at rest and able to lie flat. Examination reveals a high cervical venous column with diminished pulsation and perhaps inspiratory dilatation (Kusmaul's sign), enlargement of the liver, œdema and ascites. Pulmonary congestion and orthopnoea occur occasionally. In these cases cardiac catheterisation may reveal a raised pressure in the pulmonary artery, proving that the constriction is mainly left sided (Oglesby *et al.* 1948). The pulse is small and often paradoxical, tending to disappear during inspiration and the blood pressure is inclined to be low. Compensatory vasoconstriction may give rise to pallor, cyanosis and coldness of the skin, especially of the extremities. The heart itself does not appear to be enlarged. On palpation there may be an appreciable diastolic shock, as if the heart, filling rapidly under the influence of a high venous pressure, suddenly met the unyielding resistance of a rigid pericardium which, from a state of relaxation, was thrown abruptly into tension. On auscultation this is represented by an accentuated third heart sound or early diastolic report. Friction is absent. Auricular fibrillation occurs in about one third of cases and is the rule in patients who are over 30 years old.

Fluoroscopy reveals little cardiac pulsation, the heart shadow is normal in size in 45 per cent, slightly enlarged in 17 per cent, moderately enlarged in 32 per cent, and greatly so in 6 per cent, and has a relatively ill defined outline (Oglesby, Castleman and White 1948). Enlargement when present is usually due to the thickness of the pericardium which may measure as much as 26 mm (Freedman 1939). The shape of the heart shadow is also altered, being triangular in half the cases, with straight left and right borders and a small or absent aortic knuckle. Calcification occurs in about half the cases and is best seen in the left anterior oblique position (fig 12.67).

The electrocardiogram shows low voltage QRS complexes with flattening or slight inversion of T in all leads, representing the late stage



(a) Triangular shaped heart

(b) Calcification seen in second oblique view

Fig 12 07—Skiagrams of a case of constrictive pericarditis

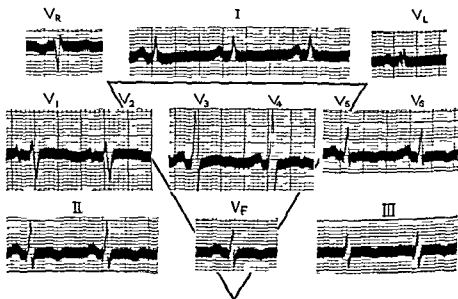


Fig 12 08—Electrocardiogram in a case of chronic constrictive pericarditis showing low voltage and flat or inverted T waves

of the pericardial T_2 pattern which in these cases is permanent (fig 12 oS)

Treatment Treatment consists of cardiac decompression achieved by surgical removal of the constricting tissue (Churchill 1929) The left side of the heart must be freed first or acute pulmonary œdema may result Removal of calcium may be difficult and time consuming but is amply rewarded The chief dangers during the operation are hæmorrhage and cardiac arrest or ventricular fibrillation The post operative course has been smoother since the advent of chemotherapy for pulmonary and pleural sepsis can now be avoided or treated effectively The frequency of positive cultures obtained from pericardial tissue removed at operation has proved that activity is no direct bar to surgical treatment but has encouraged the concomitant use of streptomycin Dramatic clinical recovery may be expected after successful pericardiectomy but the total operative mortality is about 33 per cent and another 10 per cent of cases die shortly afterwards (Sellors 1946) Persistent tuberculous activity is responsible for some of these deaths and others have been due to inexperience In expert hands the immediate surgical mortality should not exceed 10 per cent in selected cases under 50 years of age the risk in older patients is probably too great to justify operation Excellent results are obtained in 50 to 75 per cent even though the venous pressure remains somewhat elevated and the electrocardiogram shows persistent inversion of the T waves

ADHERENT PERICARDIUM

During the first quarter of this century adherent pericardium was still considered an important complication of pericarditis Extensive adhesions anchoring the heart to adjacent resistant structures were believed to add a heavy burden to ventricular systole The theory was coloured by the pathological observations of Cabot (1926) who recorded gross cardiac enlargement associated with rheumatic adherent pericarditis The clinical picture included Broadbent's sign (indrawing of the postero lateral aspect of the ribs during ventricular systole resulting from fixation of the visceral pericardium to the diaphragm) paradoxical pulse diastolic shock or rebound of the ribs fixation of the apex beat so that it failed to shift with change of posture similar fixation of the electrical axis of the heart and unexplained cardiac enlargement To cure this unhappy condition the operation of cardiolysis (Brauer 1903) was devised to free the heart of its encumbrances by dividing adhesions between it and the surrounding tissues and especially by extensive rib resection so that the heart could pull against less resistant structures In more recent years however the serious consequences of adherent pericardium have been denied and its surgical treatment is no longer favoured

Hosler and Williams (1936) failed to produce any cardiac enlargement or alteration of cardiac function by suturing the heart and pericardium to

the diaphragm in 13 dogs nor could they find a single instance of cardiac enlargement in 76 cases of adherent pericarditis in which there was not an adequate organic intracardiac cause chiefly valvular disease. Similar clinical and autopsy evidence was obtained by Armstrong (1940) in 72 cases and by Evans (Parkinson 1936) in 49 cases.

All Cabot's cases of gross cardiac enlargement with adherent pericardium were complicated by serious valve disease. Although Broadbent's sign (if not confused with indrawing of the left antero-lateral aspect of the thorax which may occur whenever the heart is grossly enlarged) and diastolic rebound of the ribs are reliable signs of extrapericardial adhesions, paradoxical pulse favours constriction and fixation of the apex beat or of the electrical axis is too variable to be of diagnostic value (France 1938).

TYPES OF PERICARDITIS BASED ON ETIOLOGY

Rheumatic pericarditis The dry form may give rise to nothing more serious than transient pericardial friction but it has an important bearing on diagnosis its advent during the course of rheumatic fever proving beyond question the presence of active carditis. More extensive pericarditis is usually associated with gross rheumatic infection so that serious carditis may be assumed. These patients are often very ill but it is not necessarily the pericarditis which makes them so. The development of cardiac dilatation and failure under these circumstances is apt to be mistaken for pericardial effusion with cardiac compression and indeed the differential diagnosis may not be easy. The position of the apex beat in relation to left border dullness the ease with which it can be felt and the presence or absence of dullness in the second left intercostal space are good guides but the electrocardiogram may be indeterminate and the interpretation of portable X-ray films difficult (the patient being too ill to move to the X-ray department). Occasionally diagnostic paracentesis may be necessary.

The electrocardiogram has failed to show any alteration of the S-T segment and T wave in at least half the cases of proved rheumatic pericarditis observed at Taplow even when subsequent necropsy has revealed a grossly thickened pericardium. No other form of pericarditis behaves like this and it is suggested that a superficial current of injury may fail to develop because of the lack of an appreciable boundary zone between the pericardium and underlying muscle owing to similar disease of both. Shortening of Q-Tc however may occur and in the absence of digitalis therapy at once distinguishes pericardial effusion from a dilated heart.

Rheumatic pericardial effusion is a clear straw coloured sterile exudate it rarely compresses the heart tends to be resorbed spontaneously without undue delay appears to respond to salicylates and is usually best left alone.

Fortunately, there are no significant after-effects for chronic constrictive pericarditis is practically never rheumatic and adherent pericardium though not uncommon is of little importance. Pericardial calcification is seen occasionally but is scanty and harmless.

Treatment is limited to relief of pain when present and to cardiac decompression in rare cases of high pressure effusion. For the former antiphlogistine is comforting but when the pain is severe morphine should be given. For the latter paracentesis is required and should be repeated when necessary. Salicylates may also help. Otherwise treatment should be directed towards the rheumatic illness as a whole.

Tuberculous pericarditis Tuberculous pericarditis may be uncommon but it is hardly rare; it affects all age groups but favours coloured races and the male sex. The infection usually spreads from mediastinal lymph glands or pleura. Effusion is the rule and if the patient survives constriction may follow. The onset is insidious and in cases with effusion a large quantity of fluid may collect before symptoms are noticed. Dyspnoea and an irritable dry cough due to pressure on the lungs and bronchi are the usual complaints. The absence of constitutional disturbances is often remarkable but continued fever, anorexia, loss of weight, night sweats and secondary anaemia may occur in the more active cases. Diagnosis depends upon the absence of rheumatic manifestations, upon the subacute or chronic course of the malady, upon the discovery of tuberculosis elsewhere and upon the results of culture and guinea pig inoculation of specimens of fluid obtained by paracentesis. The effusion is usually a clear straw coloured exudate containing lymphocytes but is sometimes blood stained. The course is prolonged, usually ranging between three and eighteen months and is often downhill with progressive emaciation, toxæmia and anaemia. Cardiac compression may become dangerous when frequent tapping adds to the patient's misery.

It is doubtful if more than 20 per cent of untreated cases with positive cultures survive and of these the majority develop chronic constrictive pericarditis subsequently, not infrequently active and constrictive stages are telescoped. The prognosis is very different when tubercle bacilli cannot be recovered from the pericardial fluid, the mortality rate being then less than 10 per cent (Harvey and Whitehill, 1937) but of course the etiological diagnosis in many of these cases is open to question and very few constrict later. Streptomycin 1 to 2 G daily for a month is giving encouraging results and may alter the outlook in proved cases.

Polyserositis Whilst tuberculosis may affect the pleura and peritoneum as well as the pericardium, the term polyserositis (Concato's disease) is usually reserved for a somewhat similar inflammatory process of unknown origin. Large effusions collect in the serous sacs, the fluid being a clear or opalescent straw coloured sterile exudate. The process is obliterative and in the pleural cavity paracentesis must be performed ever higher as the two layers of pleura become fused together in a thick dense white matting. Over the liver and spleen the greatly thickened peritoneum resembles a stout coating of sugar ice. When the pericardium is involved reabsorption of fluid is followed by total obliteration of the pericardial cavity and constriction may ensue. The course, prognosis and treatment are similar to those of tuberculous pericardial effusion.

Malignant infiltration of the pericardium When a male over 40 years of age complains of recent cough and breathlessness of insidious onset and is found to have a large pericardial effusion a malignant or tuberculous etiology is probable. If there is no fever and the fluid is blood stained the diagnostic scales tip sharply in favour of malignancy. When the pericardium is extensively invaded hemorrhagic effusion and cardiac tamponade are the rule but when it is infiltrated by a single small nodule the fluid is usually clear and straw coloured and the sac being more distensible tamponade is less frequent. The condition is invariably fatal and death never long delayed. Autopsy usually reveals a primary bronchial carcinoma.

Pyogenic pericarditis Streptococcal pneumococcal and staphylococcal infection may each give rise to pericarditis. Fever leucocytosis and toxæmia are more conspicuous than in other forms. Effusion is common and usually purulent. It is generally believed that recovery may be followed by constriction but the point is perhaps still uncertain. Streptococcal pericarditis may complicate tonsillitis erysipelas broncho pneumonia or any other streptococcal infection. It usually occurs during the acute stage of the illness and is then readily distinguished from rheumatic pericarditis but when there is an appreciable latent interval this distinction is not so easy. Pneumococcal pericarditis is usually a complication of left basal pneumonia organisms gaining access to the pericardium by direct spread from the pleura. Staphylococcal pericarditis may complicate myocardial abscess from staphylococcal septicæmia.

The course and prognosis of pyogenic pericarditis have been radically altered by chemotherapy. Penicillin is more effective than the sulphonamides and should be given in doses of 30 000 units intramuscularly every three hours day and night for seven to ten days. About 10 to 20 ml of a solution containing 1 000 units of penicillin per ml should also be injected into the pericardial sac after paracentesis. Surgical drainage is only necessary when there is frank suppuration. With such treatment initial recovery is the rule but the ultimate outcome is uncertain. The low mortality rate may result in an increased incidence of chronic constrictive pericarditis on the other hand the prevention of frank suppuration may have the opposite effect. The few cases so far followed up by the author have not constricted.

Benign pericarditis of uncertain etiology Most physicians are familiar with a benign form of pericarditis which may complicate upper respiratory tract infections as described by Logue and Wendkos (1948) or which may arise spontaneously. Effusion when present is sterile and complete recovery takes place without treatment. Some of these are probably pyogenic in origin and some are undoubtedly tuberculous, as the subsequent history may show. It is therefore wise to treat all such cases with penicillin or streptomycin according to whether the clinical features correspond more closely to the pyogenic or tuberculous variety.

Hæmopericardium and traumatic pericarditis Hæmorrhage into the pericardial sac may be caused by stab or gun shot wounds by rupture of a

syphilitic or dissecting aneurysm of the aorta or by perforation of a myocardial infarct or ventricular aneurysm. Wounds of the heart are not necessarily fatal and if the patient survives the initial insult relief of cardiac tamponade and surgical repair may be life saving. Rupture of the heart or aorta into the pericardium is always fatal but not necessarily immediately. A patient with a perforated infarct for example may live as long as ten days.

Severe blows or crush injuries to the chest or blast may cause myocardial bruising and pericardial ecchymoses. Transient pericardial friction and characteristic electrocardiographic changes usually provide evidence of the lesion. If there is no damage to the superficial coronary arteries complete recovery is the rule.

An interesting form of traumatic pericarditis may be due to pericardial foreign body (usually a metallic fragment) or to a foreign body lying close to the pericardium. In these cases recurrent attacks of pericarditis with clear sterile effusion may occur at any time up to four months after the original injury. The interval between attacks is usually two to six weeks during which the patient seems perfectly well. The attacks themselves which last about a week tend to be severe with fever rapid effusion cardiac tamponade and considerable pain and distress. Of seven cases reported by Wood (1945) however none died. If the foreign body is easily accessible it is best removed in a quiescent period if not it may be safer to leave it *in situ*.

Uræmic pericarditis Pericardial friction is not uncommonly heard in patients dying of uræmia. Symptoms are rare effusion absent and electrocardiographic changes minimal. At autopsy needle like crystals of urea may be found massed in the pericardium.

Pericarditis secondary to myocardial infarction Acute myocardial infarction may give rise to a local (60 per cent) or general (15 per cent) pericardial reaction and perforation may lead to hæmopericardium. Local pericarditis is limited to the surface area of the infarct gives rise to no symptoms and does not interfere with the electrocardiographic pattern of the underlying lesion. A fleeting friction rub may be heard if the infarct is anterior.

General pericarditis is less



Fig. 12.07—Pericardial effusion of three years' duration in a case of extreme essential hypertension.

Intra-pericardial pressure was 9 cm. of saline and the venous pressure was raised. The patient was able to lie flat or to be tilted head down without breathlessness. After paracentesis orthopnoea and paroxysmal cardiac dyspnoea developed within a few days.

common but more important for it may cause additional pain, allows anterior friction to be associated with posterior infarction (Stewart and Turner 1938) influences the electrocardiographic pattern and may even give rise to effusion

Hydroperticardium Hydroperticardium associated with congestive failure is never conspicuous and is of no clinical significance. As a complication of myxœdema pericardial effusion is not uncommon. An interesting though rare variety of hydroperticardium is occasionally encountered in cases of malignant hypertension (fig. 12.09). Relatively high pressure may develop in the sac and by hampering cardiac filling may prevent pulmonary congestion. Such cases may remain remarkably free from symptoms if the effusion is tapped. orthopnoea and paroxysmal cardiac dyspnoea may develop.

REFERENCES

- Armstrong T G (1940) Adherent pericardium constrictive and non constrictive *Lancet* ii 475
- Brauer L (1903-4) Die kardiolyse und ihre indicationen *Arch f Chir* 71 258
- Broadbent W H and Broadbent J F H (1897) Heart disease with special reference to prognosis and treatment New York
- Burchell H B Barnes A R and Mann F C (1939) The electrocardiographic picture of experimental localised pericarditis *Amer Heart J* 18 133
- Cabot R C (1926) Facts on the heart Philadelphia
- Capps J A (1932) An experimental and clinical study of pain in the pleura pericardium and peritoneum New York
- Chevers N (1842) Observations on the disease of the orifice and valves of the aorta *Cut Hosp Rep* 7 387
- Churchill F D (1929) Decortication of the heart (delorme) for adhesive pericarditis *Arch Surg* 19 1457
- Fwart W (1896) Practical aids in the diagnosis of pericardial effusion in connection with the question as to surgical treatment *Brit med J* i 717
- France R (1938) The use of the electrocardiogram in the diagnosis of adhesive pericardio mediastinitis *Bull Johns Hopk Hosp* 63 104
- Freedman E (1939) Inflammatory diseases of pericardium *Amer J Roentgenol* 42 38
- Harvey A M and Whitehill M R (1937) Tuberculous pericarditis *Medicine* 16 45
- Hosler R M and Williams J E (1936) Study of cardio pericardial adhesions *J thorac Surg* 5 629
- Kisch B Nahum L H and Huff H E (1940) The predominance of surface over deep cardiac injury in producing changes in the electrocardiogram *Amer Heart J* 20 174
- Logue R B and Wendkos M H (1948) Acute pericarditis of benign type *Ibid* 36 587
- Lower R (1669) Tractatus de Corde I eyden ed 1728 Quoted by Major R H (1932) Classic descriptions of disease Springfield Illinois
- Oglesby P Castleman B and White P D (1948) Chronic constrictive pericarditis *Amer J med Sc* 216 561

- Parkinson J (1936) Enlargement of the heart *Lancet* **i** 1341
- Pick F (1896) Über chronische unter dem Bilde der Lebercirrhose verlaufende Perikarditis (perikarditische Pseudolebercirrhose) meiste Bemerkungen über die Zuckergussleber (Curschmann) *Ztschr F klin med* **29** 385
- Porte D and Pardee H E B (1929) Occurrence of coronary T wave in rheumatic pericarditis *Amer Heart J* **4** 584
- Rotch T M (1878) Absence of resonance in the fifth right interspace diagnostic of pericardial effusion *Boston med & surg J* **99** 389 421
- Sellors T H (1946) Constrictive pericarditis *Brit J Surg* **33** 215
- Stewart H J Crane N F and Deitrick J E (1938) Studies of the circulation in pericardial effusion *Amer Heart J* **16** 189 — and Heuer G J (1939) Chronic constrictive pericarditis *Arch intern Med* **63** 504 — and Turner K B (1938) A note on pericardial involvement in coronary thrombosis *Amer Heart J* **15** 232
- White P D (1935) Chronic constrictive pericarditis (Pick's disease) treated by pericardial resection *Lancet* **ii** 597
- Wood P H (1937) Electrocardiographic changes of a T pattern in pericardial lesions and in stab wounds of the heart *Lancet* **ii** 796 — (1945) War wounds of the heart *Proc Conf Army Phys Rome* **23**

SYPHILITIC AORTITIS

SYPHILITIC inflammation of the aorta is clinically unrecognisable unless it results in fusiform dilatation, saccular aneurysm, aortic incompetence, angina pectoris or possibly heart block. It is true that many museums contain a specimen of syphilitic myocarditis and even of syphilitic endocarditis but these are oddities. The various manifestations of syphilitic aortitis commonly appear from ten to thirty years after primary infection, usually between the ages of 40 and 60, account for about 5 per cent of all cases of organic heart disease in Britain and are approximately five times more common in men than in women.

There can be little doubt that the disease is becoming less frequent and will become rare. This is of course the result of educating the public in venereology and in the improved treatment of early syphilis. Thompson, Comeau and White (1939) found cardiovascular syphilis had developed clinically in 10 per cent of 241 patients known to have had syphilis fifteen to twenty five years previously; all had been inadequately treated by 1939 standards. Uncomplicated aortitis, rarely recognised except at necropsy, is undoubtedly much more frequent but its exact incidence is difficult to assess, published figures depending on the criteria upon which the diagnosis rests. Aortic incompetence is about twice as common as aneurysm.

Whilst the clinical features may be diagnostic of a syphilitic etiology, the latter may be confirmed by a history of syphilis, by signs of syphilis in other systems (particularly neurosyphilis), by a positive Wassermann or Kahn reaction in the blood in about 85 per cent of cases, and by a persistently raised erythrocyte sedimentation rate.

Congenital syphilis does not cause aortitis (McCulloch 1930) although spirochaetes may be present in the aorta; there is practically no tissue reaction.

Pathology. The initial lesion occurs in the adventitia and consists of syphilitic endarteritis of the vasa visorum and of focal granulomatous tissue. Although the inflammation spreads deeply into the media, atrophy and necrosis of muscular and elastic fibres are partly due to ischaemia. The damage is patchy and is repaired by fibrous tissue, the cross section of the aortic wall being correspondingly thinned at such points. These medial scars are indicated on the inner surface of the vessel by depressions of the intima which presents a pock-marked appearance. Secondary atherosclerosis with extensive calcification is common.

ANEURYSM

Sooner or later the diseased media yields to the force of the blood pressure either generally or at its weakest point and a fusiform or saccular aneurysm results. A fusiform aneurysm is little more than an exaggeration of the inevitable dilatation of a syphilitic aorta and has no greater consequences. It is usually associated with aortic incompetence but it may be seen radiologically when still uncomplicated; it then affords the only acceptable clinical evidence of relatively early syphilitic aortitis. The diagnosis should be confirmed by a history of syphilis or by a positive Wassermann or Kahn reaction in the blood; however, for fusiform aneurysm may result from non-specific medial necrosis with or without dissection and slight dilatation of the aorta may be due to atherosclerosis and hypertension. A ringing or amphoric second heart sound at the base of the heart may denote dilatation of the ascending aorta but does not indicate its cause. Again a suspicious aortic second sound must be disregarded if the ascending aorta is seen to be normal in size and shape.



FIG. 1301.—Saccular aneurysm of the ascending aorta

The syphilitic aneurysm proper is saccular (fig 1301) and may occur in any part of the thoracic aorta, particularly in the arch. Aneurysm of the abdominal aorta is rare and is more often atheromatous in origin.

Incidence. The sex incidence of saccular aneurysm is male to an even greater degree than other varieties of late syphilis, the sex ratio approaching 10:1 in their favour (White 1937). This is probably due to the greater stresses imposed on the aorta in men. It is significant that saccular aneurysm rarely develops when there is aortic incompetence, occurring in little over 10 per cent of such cases (Welty 1939) and that symptoms due to pressure from an aneurysm may be relieved by an artificial arterio-venous shunt conditions which reduce the mean aortic pressure.

ANEURYSM OF THE ASCENDING AORTA

Aneurysm of the ascending aorta usually causes visible pulsation or a conspicuous pulsating tumour to the right or left of the sternum or in the suprasternal notch. Symptoms may be absent or there may be sternal or

costal pain from pressure erosion. Pulsation may be expansile and may be accompanied by a systolic thrill. On auscultation a loud systolic bruit is usually heard. When invisible an anterior aneurysm may yet be detected by percussing a band of parasternal dullness.

Partial obstruction of the superior vena cava is a common complication and gives rise to a high venous pressure in the head and neck while the

right auricular pressure remains normal. The distended jugular veins do not pulsate unless the obstruction is slight and may be readily overlooked. A visible collateral venous circulation does not necessarily develop presumably because the block is incomplete so that a fair blood flow is maintained under the high head of pressure. Puffiness or œdema of the head may occur and in one case of the author's gravitated to the legs in the erect posture. The diagnosis may be proved by means of angiocardiology (fig 13 02) or by passing a venous catheter and noting the sudden fall in pressure as the tip slips through the obstruction.



Fig 13 0 —Angiocardiogram showing partial superior vena cava obstruction due to an aneurysm

(B r v f D F (d)

Aneurysm of the ascending aorta may rupture into the pericardial or pleural cavities

into the pulmonary artery or into the right auricle

ANEURYSM OF THE ARCH

The symptoms and signs of an aneurysm in this situation are determined by pressure on surrounding structures. Practically any structure in or close to the superior mediastinum may be compressed according to the size and position of the aneurysm. Thus pressure on one or other subclavian artery may lead to significant differences in the pulse and blood pressure in the two arms; a rare complication of this is clubbing of the fingers on the affected side. Pressure on the left bronchus causes collapse of the left lung which may be complete or partial, the upper lobe being involved more often than the lower. Inflammatory changes may occur distal to the obstruction, particularly bronchiectasis and pulmonary tuberculosis. The left bronchus may be depressed with each pulsation of the aneurysm; the resulting downward pull on the trachea during systole may be readily

detected at the cricoid cartilage and is known as a tracheal tug. It is best elicited by standing behind the patient who should be seated and applying steady upward pressure on the cricoid cartilage with the tip of one fore finger. Pressure on the trachea itself may give rise to an irritating cough to stridor or to considerable respiratory obstruction. Pressure on the left recurrent laryngeal nerve to a brassy cough and paralysis of the left vocal chord. Pressure on the œsophagus to dysphagia. The phrenic nerve usually escapes as it lies superficially but the left sympathetic chain may be compressed with the production of Horner's syndrome—homolateral contraction of the pupil and partial ptosis. Severe radiating pains may be caused by pressure on nerve roots and the spine may be eroded.

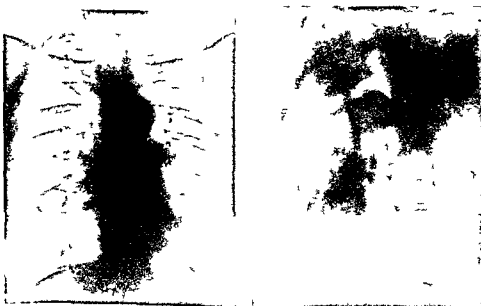
ANEURYSM OF THE ABDOMINAL AORTA

Of 1459 aneurysms of the aorta collected from the literature by Ungar and Poppel (1936) only 136 or less than 10 per cent were below the diaphragm.

An abdominal aneurysm usually presents as a pulsating tumour in the epigastrium over which a systolic thrill and murmur may sometimes be detected. Root pain associated with vertebral erosion is not uncommon or pain may be local. A relatively common clinical error is to mistake a normal aorta projected forwards by lordosis in thin subjects for an aneurysm.

RADIOLOGICAL DIAGNOSIS

Although the existence and site of a syphilitic aneurysm may be diag-



(a) Antero-posterior view

(b) Second oblique posterior view

Fig. 1303—Skiagram showing several aneurysms of the aortic arch.

(By courtesy of J. S. Park.)



Fig 13 04—Skilogram showing erosion of the bodies of several dorsal vertebrae as the result of pressure from an aneurysm



Fig 13 05 (a)



Fig 13 05 (b)



Fig 13 05 (c)

Fig 13 05—Angiocardiogram outlining a normal pulmonary artery (a) and aorta (b) in a case of mediastinal tumour (c)

nosed clinically with a fair degree of accuracy radiological confirmation should always be obtained. Aneurysm may be distinguished from other rounded shadows in the vicinity of the aorta by having four characteristic features (1) it is intimately connected with the aorta (fig 13 03) (2) it becomes opaque with the aorta in angiocardiograms (3) it pulsates unless it is thrombosed (4) some part of its wall may be calcified (fig 13 03a). Erosion of the bodies of several vertebræ (fig 13 04) and compression of the trachea bronchus or œsophagus may sometimes be seen. Confusion may arise however when a mediastinal tumour exhibits transmitted pulsation. Angiocardiography is advised in all doubtful cases (fig 13 05).

COURSE

Many aneurysms remain silent and are discovered accidentally by radiography others cause much suffering. One of the worst features is the severe pain produced by pressure on bone especially the root pain associated with vertebral erosion. This may last for months and be very resistant to treatment.

The prognosis varies greatly but the average duration of life is little more than eighteen months from the onset of symptoms (Colt 1926-27). Cases have been reported however which have survived for fifteen to thirty years (Kauntze 1947). The chief dangers are infection of the lungs distal to bronchial compression and rupture. Aortic aneurysm may rupture into the pericardium into the pulmonary artery into the trachea or bronchus into the œsophagus or into the pleura giving rise to hæmopericardium with cardiac compression to acute right ventricular failure with signs and symptoms of an aorto pulmonary shunt to dramatic hæmoptysis to hæmatemesis or to hæmothorax respectively usually with fatal results.

SPECIAL TREATMENT

The object of treatment apart from anti syphilitic measures is to promote thrombosis and calcification in the aneurysmal sac or to protect it by means of external fibrosis in order to prevent rupture or further expansion.

Bed rest is necessary at first while routine anti syphilitic treatment is given (page 367). During this period a course of calcium lactate 10 grains (0.6 G) t d s with vitamin D may be added to promote calcification in the wall of the aneurysm.

If pain is not relieved by these measures surgical interference may be considered. The old operation of inserting a wire into the sac in order to induce thrombosis is unsatisfactory the risk is considerable and effective clotting cannot be guaranteed. Babcock's operation—the creation of an arterio venous communication between the carotid and jugular vessels (Babcock 1926 1932)—reduces the mean aortic pressure and may relieve

pain (Ranson 1947) The most promising surgical method however, is to wrap the aneurysm in polythene cellophane this causes an intense fibroblastic reaction which protects the sac from without prevents further expansion and relieves pain (Poppe 1948)

AORTIC INCOMPETENCE

Pathology Weakening of the meso-aorta in the region of the aortic valve leads to dilatation of the aortic ring and to separation of the cusps at their commissures so that the valve becomes incompetent Granulomatous tissue may also drive a wedge between the junctions of the cusps The cusps themselves become rolled and thickened at their free margins and present a dwarfed stunted appearance There is no stenosis and calcification is absent unless there is much secondary atherosclerosis Owing to the site of the lesion which is necessarily at the root of the aorta the mouths of the coronary vessels are often scarred and narrowed indeed they may be almost occluded Ischaemic fibrosis of the myocardium results

Incidence Syphilis accounts for about one third of all cases of aortic valve disease and for about one half of those in subjects between the ages of 40 and 60 (Cowan and Ritchie 1935)

The sex ratio in syphilitic aortic incompetence is about 3 : 1 in favour of men (Campbell 1932) and is thus less remarkable than in aneurysm

Clinical features Syphilitic aortic incompetence has all the features of aortic incompetence in general (page 295) and some special characteristics of its own Only the latter will be considered here

1 The history of symptoms or of the discovery of the lesion is relatively recent usually a matter of weeks or months and rarely more than a year or two

2 Angina pectoris is common occurring sooner or later in about half the cases

3 The aortic incompetence is pure there being no stenosis (unless there is secondary atherosclerosis) and therefore no systolic thrill

4 The incompetence is usually very free so that peripheral vascular manifestations are marked

5 The murmur is apt to be to and fro replacing the heart sounds at the base and is often heard better at the aortic area than down the left border of the sternum owing to dilatation of the ascending aorta

6 A basal diastolic thrill is suggestive being very rare in other types of aortic valve disease except when a cusp is perforated or ruptured

7 Skiagrams may reveal differences in the diameter of the aorta at different points or fusiform aneurysm (fig 13.06) but an associated saccular aneurysm is rare Irregularity of the lumen may be seen well in angiograms

8 Fluoroscopy shows less aortic pulsation than in equivalent rheumatic cases owing to the loss of elasticity

9 Calcification of the aortic valve is against syphilis but may occur when there is secondary atherosclerosis calcification of the ascending aorta is in favour of syphilis

10 The electrocardiogram more often shows bundle branch block or various degrees of heart block than it does in rheumatic aortic incompetence but no more so than in calcific aortic stenosis The changes are attributable to chronic fibrosis rarely to gummatous lesions



(a) Antero posterior view



(b) Left anter or oblique position

Fig 13 06—Skigram of a case of syphilitic aortic incompetence showing fusiform dilatation of the ascending aorta and gross enlargement of the left ventricle

The diagnosis may be confirmed by obtaining other evidence of syphilis as previously described The FSR is more often raised in syphilitic aortic incompetence than in any other variety of late syphilis including aneurysm and is probably as good a guide to activity as the Wassermann reaction

Course The prognosis is bad the average duration of life being about two years (Campbell 1932) Left ventricular failure develops sooner or later in many cases and congestive heart failure follows The downhill course differs from that of other forms of aortic incompetence in its rapidity in the frequency of angina pectoris and in the relatively high proportion of sudden deaths (Munck 1946)

ANGINA PECTORIS

Pathology It is often said that aortic valve disease may cause angina pectoris. Whilst this is true the statement needs amplification. Angina is a common complication of all forms of aortic stenosis and of *syphilitic* aortic incompetence but not of other varieties. Rheumatic aortic incompetence for example must be gross to cause angina and rarely does so. The explanation is to be found in the physiology of the coronary circulation. During systole ventricular contraction prevents blood flowing through coronary vessels which penetrate left ventricular muscle and large arteries on the surface dilate to form an elastic reservoir which in recoil during diastole acts as an accessory pump forcing the blood onwards. The higher the systolic pressure the greater the elastic reservoir, provided the coronary arteries are healthy. During ventricular relaxation blood is able to flow through vessels penetrating muscle being propelled by the aortic diastolic pressure and by the recoil of the elastic reservoir just mentioned. Thus the coronary flow depends upon both systolic and diastolic pressures i.e. upon the mean pressure.

Now in aortic stenosis the mean blood pressure is often low but in aortic incompetence although the diastolic pressure may be 40 or 50 mm of mercury the systolic pressure is commonly raised and the mean pressure adequate. Syphilitic aortic incompetence causes angina pectoris because there is associated stenosis of the mouths of the coronary arteries. If syphilitic aortitis produces sufficient damage in the region of the aortic cusps to cause aortic incompetence it is unusual for the mouths of the coronary vessels to remain unscathed. Conversely if the mouths of the coronary vessels are so stenosed as to cause angina pectoris it is practically impossible for the root of the aorta to remain healthy. Thus syphilitic angina is rare without aortic incompetence.

Clinical features Syphilitic angina has certain characteristics which help to distinguish it from other types. (1) the attacks tend to be of longer duration (2) they are more often nocturnal although the ordinary relationship to effort holds good (3) they are less often relieved by trinitrin. Coronary thrombosis is a rare complication because ischaemia is due to stenosis of the coronary ostia and not to changes in the coronary vessels themselves. Myocardial infarction however may occur without coronary thrombosis especially when the effect of gross occlusion of the mouths of the coronary vessels is exaggerated by a drop in blood pressure from some other cause such as surgical shock. Ischaemia of the least nourished part of the myocardium may then be so pronounced as to cause necrosis even so cardiac infarction is uncommon.

Course The prognosis is bad patients rarely living more than two years. Status anginosus may develop before the end or nocturnal angina may prove troublesome. When heart failure develops angina may disappear it is not clear why this should be so but it may depend upon changes in tissue metabolism.

Special treatment The ordinary method of treating angina pectoris are applicable to the syphilitic type although the results are poor. Anti syphilitic measures should not be withheld but arsenic must be given with special care. Bed rest is particularly important during the first six weeks of treatment and is essential during the first course of penicillin or arsenic.

HEART BLOCK

True syphilitic heart block is very rare and depends upon interruption of the bundle of His by gummatous tissue. On the other hand heart block resulting from interference with the conducting tissue by ischæmic fibrosis due to stenosis of the coronary ostia is not uncommon and like angina pectoris and for the same reason is nearly always associated with aortic incompetence. The former responds to iodine therapy the latter of course does not.

TREATMENT OF SYPHILITIC AORTITIS

Syphilitic aortitis should be fully treated with anti syphilitic drug whether uncomplicated or otherwise. Clearly past damage cannot be repaired but further activity can be prevented. The objective is to secure a negative Wassermann reaction and a normal sedimentation rate.

The patient should be put to bed for six weeks during which period he should receive potassium iodide 10 grains (0.6 G) t d s preferably with liq. hydrarg. perchlor. 60 minims (4 ml). Gummatous tissue resolves quickly with this treatment and the danger of a severe Herxheimer reaction is lessened. After three to four weeks of treatment with iodides a minimum of 2.4 million units of penicillin may be given over a period of 10 days in divided doses of 30,000 units three hourly. The course may well be modified in the light of experience but the total dose should not be less. Reactions are rare whether the initial dose is 1,000 or 100,000 units (Moore 1947).

Shortly after completing the penicillin course the patient may be allowed up. Iodide and mercury are then discontinued and treatment with bismuth should be instituted. Intramuscular injections of 0.1 G of bismostab twice weekly for six weeks followed by 0.2 G weekly for the next six weeks are advised.

The patient is then put back to bed and arsenic is given intravenously beginning with 0.15 G of N A B and continuing with 0.3 G, 0.45 G and 0.6 G at weekly intervals the last dose being repeated until a total of 4.5 G is reached. If there is no untoward reaction after the first two 0.6 G doses of N A B the patient may be allowed up again. The development or aggravation of angina pectoris or heart failure are major dangers but both are unlikely and when they do occur cannot always be attributed to the treatment. Nevertheless arsenic should be abandoned should they occur.

The regime described constitutes one complete course of treatment and lasts six months. The situation should then be reviewed, particular attention being paid to the Wassermann reaction and the C S R. If both are normal treatment may be discontinued for six months but if either suggest persistent activity the whole procedure with certain modifications should be repeated without delay. Bed rest is no longer necessary unless congestive failure, angina pectoris or other complications demands it and bismuth may be given in doses of 0.2 G. and N A B in doses of 0.6 G. from the first injection.

Two complete courses are strongly advised in all cases with an interval of 12 months or consecutively. A third or fourth course should be given without hesitation if there is any further evidence of activity.

It is repeated for emphasis that neither aneurysm, angina pectoris nor heart failure contraindicate penicillin or arsenicals provided small doses are employed initially for the Herxheimer reaction is rare. This phenomenon consists of a local tissue reaction which may cause swelling and occlusion of the coronary ostia with disastrous results even where there is no dramatic ill effect the patient may slip downhill with increasing angina or heart failure.

Statistics have shown that the effect of full anti-syphilitic measures before the introduction of penicillin was to improve the average life expectancy from eighteen months to two years (Padgett and Moore 1935). It may be argued that the increased care and enforced rest which are a necessary corollary of this form of treatment might also improve the prognosis by a similar amount and there is something to be said for the view that there is little point in attempting to extirpate the spirochæte once heart failure or angina pectoris has developed for it is then probably too late. However in the light of present knowledge it is wiser to give full treatment to all cases. Whether penicillin therapy will further increase life expectancy remains to be seen.

Complications should be treated as they arise and by the customary methods. It is wise to avoid arsenic and probably penicillin in cases of heart failure or angina decubitus until these have been satisfactorily controlled.

REFERENCES

- Babcock W W (1926) New treatment of thoracic aneurysm. *Ann clin Med* 4 933 — (1932) Newer surgical methods of treating diseases of vascular system. *Amer J Surg* 16 401.
- Campbell M (1932) A note on aortic valvular disease. *Brit med J* 1 20.
- Colt G H (1926-7) Clinical duration of saccular aortic aneurysm in British born subjects. *Quart J Med* 20 331.
- Cowan J and Ritchie W T (1935) Diseases of the heart. London p 241.
- Kauntze R (1947) Unusual longevity in aneurysm of the thoracic aorta. *Brit Heart J* 9 96.

- McCulloch H (1930) Congenital syphilis as a cause of heart disease *Amer Heart J* 6 136
- Moore J E (1947) Discussion on the treatment of syphilis with penicillin *Proc Roy Soc Med* 40 811
- Munck W (1946) The pathological anatomy of sudden death *Acta Path et Microbiologica Scandinavica* 23 107
- Padget P and Moore J E (1935) The results of treatment in cardiovascular syphilis *Amer Heart J* 10 1017
- Poppe J K (1948) Cellophane treatment of syphilitic aneurysms with report of results in six cases *Ibid* 36 252
- Ranson F T (1947) Babcock's operation for thoracic aneurysm *Brit med J* 11 69
- Thompson W P Comeau W J and White P D (1939) The role of the treatment of syphilis in the prevention of cardiovascular involvement *Amer Heart J* 17 286
- Ungar A S and Poppel M H (1936) Aneurysm of abdominal aorta report of case exhibiting auricular calcifications *Amer J Roentgenol* 36 5-3
- Welty J W (1939) Necropsy survey of cardiovascular syphilis with particular reference to its decreasing incidence *Amer J med Sc* 197 78
- White P D (1937) Heart disease New York

CHAPTER XIV

ISCHÆMIC HEART DISEASE

DEFINITION

OCCLUSIVE disease of the coronary arteries of sufficient degree to prevent the coronary circulation meeting the physiological demands of the heart is best described as ischæmic heart disease. It is characterised clinically by angina pectoris and cardiac infarction pathologically by occlusive coronary atherosclerosis with or without thrombosis and by focal ischæmic myocardial necrosis and fibrosis.

INCIDENCE

Occlusive coronary atherosclerosis is responsible for about 20 to 25 per cent of all types of organic heart disease and for about 80 per cent of all sudden cardiac deaths (Munck 1946) moreover it appears to be increasing rapidly thus the number of cases dying from coronary disease in England per million persons living was 48 in 1926 148 in 1930 and 473 in 1939 (Cassidy 1946). The increasing age of the population is no doubt partly responsible thus the citizens of ancient Rome in their halcyon days had an average life span of twenty to thirty years and the following table shows the increased average life span in the U.S.A. from 1879 to 1944 (Master 1947).

1879-1899	34 years
1911-1912	46.63 years
1919-1920	51.14 years
1930	57.36 years
1944	64.40 years

These remarkable figures are chiefly due to the successful war against infections and parasitic diseases and to the saving of life by surgical means. Another factor that must be taken into account is the attitude of medical practitioners who in 1926 had scarcely heard of coronary thrombosis whereas now they are apt to diagnose it more frequently than it exists. It should be remembered that coronary thrombosis was not recognised as a clinical entity until its classic description by Herrick in 1912 and was not widely appreciated in England until popularised by McNee in 1925.

See Of Heberden's 100 cases of angina only three were women (Heberden 1802). Most investigators give the general sex ratio as 4 : 1 in favour of men but under the age of 50 it is 8 : 1 (Hedley 1939) and under the age of 40 male predominance is overwhelming in fact angina in women under 40 is nearly always due to some other etiological agent such as hyper

tension aortic stenosis syphilitic aortitis anaemia myxœdema diabetes mellitus xanthomatosis or paroxysmal rhythm change Between the ages of 60 and 70 however about one third of the cases are women and over the age of 70 the sex incidence is equal (Gordon Bland and White 1939)

Age Of 1 000 cases seen personally by Cassidy (1946) 70 per cent were between 50 and 70 years of age of the men 14.6 per cent were between 40 and 50 3.2 per cent between 30 and 40 and 0.25 per cent were under 30 These figures are in harmony with common experience except perhaps with regard to the incidence in young men for many such cases were seen in the Services during the second world war (Newman 1946 Poe 1947) The peak age of death is 60 (Hedley 1939)

Habits and occupation There is a general impression that the incidence of ischæmic heart disease is particularly high amongst professional men and is related to the stress of modern urban life There is said to be little to support this view (Master 1947) but some figures published by Hedley (1939) are interesting

<i>Occupation</i>	<i>Deaths from coronary occlusion per 100 000</i>
Professional	154
Managers and officials	140
Clerks and salesmen	128
Skilled and unskilled workers	107

The author however, ascribed the difference in these figures to more accurate certification in those earning larger incomes

Nevertheless the obstinate belief that angina pectoris is a doctor's disease persists and has at last been justified by startling figures published by Ryle and Russell (1949) These workers who are especially well qualified to sift and present evidence of the kind required divided the social strata of England and Wales into five classes and found that the standardised mortality ratio (S M R) from ischæmic heart disease in social class I (professional workers) was twice that in social class III (skilled artisans) and three times that in class V (unskilled workers) Their table giving the actual occupations with the four highest and four lowest standard mortality ratios ends the debate on this previously vexed question and once again emphasises the fact that experienced opinion should not be too readily cast aside because of ill founded statistical evidence to the contrary Physicians and surgeons head the list with an S M R of 368 proprietors of wholesale business came second with an S M R of 235 the legal profession third (227) and the Church fourth (218) At the other extreme we have workers in chemical processes with an S M R of only 20 agricultural labourers 32 stone miners and quarriers 38 and coal miners engaged in other work 40

There is no evidence that either alcohol (Wilens 1947) or tobacco (Cassidy 1946) is responsible for the high male incidence or has any permanent influence on the course of the disease although heavy drinking or smoking may provoke attacks of angina

PATHOGENESIS

Ischæmic heart disease is due to occlusive coronary atherosclerosis with or without secondary subintimal hæmorrhage or thrombosis. Angina pectoris caused by syphilitic aortitis, aortic stenosis, severe anæmia, paroxysmal



Fig. 1401 (1).—Skiazgram of the heart in a case of occlusive coronary atherosclerosis: the coronary vessels have been injected with a radio opaque gel.

tachycardia and the like, and coronary occlusion resulting from angitis, embolism, trauma, dissecting aneurysm and other rarities are considered elsewhere.

The cause of human atheroma remains unknown, despite a great deal of

work on the subject (Cowdry 1933) Lipoid substances accumulate in the intima of the aorta and larger arteries in a patchy irregular fashion causing a variable degree of pressure atrophy of the underlying media and some times encroaching on the lumen of the vessel (fig 14 01)



Fig 14 01 (b)—Normal control for comparison

The lipoid nature of the deposits their relatively frequent association with diabetes mellitus and hypercholesterolaemia and their easy reproduction in rabbits by cholesterol feeding (Leary 1934) suggest some relationship to fat metabolism The manner in which lipoid laden macrophages (lipophages) swept to the side of the blood stream owing to their

light weight, may penetrate the intima has been recently described by Gordon (1947) hypertension facilitates the process

The degree of narrowing of an atherosclerotic coronary artery cannot be accurately assessed by its appearance at necropsy for in life the blood pressure tends to iron out the excrescences and maintain a smooth intimal surface and full lumen (Harrison and Wood 1949)

Although the intima and early atheroma are avascular advanced lesions develop a blood supply from the vasa vasorum (Leary 1938) and sub intimal hæmorrhage may then occur causing sudden occlusion of the vessel (Paterson 1936 1939 1941 Wartman 1938) such accidents however are rare and account for only about 1 per cent of cases of sudden coronary occlusion

Secondary calcification of advanced atherosclerosis may convert the coronary arteries into bony tubes as in the classical example of John Hunter (1796) Erosion or ulceration of atheromatous lesions forms an excellent nidus for secondary thrombosis This is the common cause of acute coronary obstruction Organisation of such thrombi leads to microscopical appearances similar to atherosclerotic lesions indeed it has even been suggested that atheroma may represent nothing more than intravascular clotting (Duguid 1946)

ANGINA PECTORIS

Physiology Angina pectoris and its close relative the pain of intermittent claudication are believed to be due to certain metabolites that are formed in ischæmic working muscle (Lewis 1934) Whatever the precise explanation for the development of pain there can be no doubt that attacks depend upon relative myocardial ischæmia an idea first enunciated by Parry (1799) The term *angina pectoris* is customarily applied to transient pain only and refers to ischæmic attacks provoked by temporary stress during which the metabolic demands of the myocardium are beyond the capacity of the coronary circulation

Such a situation may arise during effort (1) if the coronary vessels are more or less occluded either at their mouths as in syphilitic aortitis (page 366) or during their course as in atherosclerosis various forms of angitis and embolism (2) if the coronary flow is diminished by other means such as aortic stenosis (page 300) gross aortic incompetence or congenital anomaly (3) if the blood itself carries insufficient available oxygen as in anæmia or at high altitudes or (4) if the regular work of the heart is increased by such conditions as hypertension aortic valve disease thyrotoxicosis or anæmia

Although only angina pectoris resulting from coronary atherosclerosis concerns us here the other factors mentioned often play a contributory role thus anæmia may precipitate angina in a case of previously silent coronary disease not only because of the limited oxygen supply but also

because the work of the heart is increased in order to maintain a high cardiac output. Hypertension is particularly important in so far as it increases the work of the heart and contributes to the development of atheroma; on the other hand it tends to iron out the plaques and so may prevent coronary narrowing. In fact most cases of hypertensive heart disease have dilated coronary arteries (fig 15 09) (Harrison and Wood 1949). Clinically although more than half of all cases of ischæmic heart disease have blood pressures above 160/100 mm Hg (Cassidy 1946) systolic pressures over 200 mm Hg are rare (Riseman and Brown 1937).

CLINICAL FEATURES

Angina is but a symptom and may be distinguished from other pains in the upper half of the body by a careful analysis of its qualities and behaviour.

Site The pain is central mid sternal and tends to radiate bilaterally across or round the chest into the sides of the neck and jaws or even into the face or nose into the shoulders and down the inner or outer sides of the arms sometimes as far as the little fingers or thumbs occasionally through to the back between the shoulder blades (fig 14 02). This full distribution was experienced by John Hunter (1796). It is not situated in the left breast area although it may be rather to the left of the sternum than in the mid line; a localised left inframmary pain is never angina. Radiation may be unilateral and it is true that the left side then suffers more often than the right but it must not be thought that spread down the left arm is either especially typical or diagnostic; for bilateral spread is more typical and many other pains may radiate down the left arm including left inframmary pain. Although centrifugal spread is the rule radiation is occasionally centripetal the pain starting in the wrists upper arms or face and spreading thence to the chest. Pain may even be confined to one of the points of radiation e.g. to the face back or wrist not being felt in the front of the chest at all.

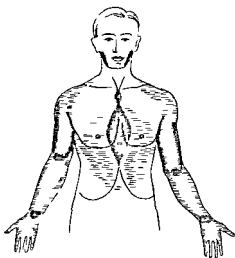


Fig 14 02—Diagram illustrating radiation of pain in ischæmic heart disease

the right but it must not be thought that spread down the left arm is either especially typical or diagnostic; for bilateral spread is more typical and many other pains may radiate down the left arm including left inframmary pain. Although centrifugal spread is the rule radiation is occasionally centripetal the pain starting in the wrists upper arms or face and spreading thence to the chest. Pain may even be confined to one of the points of radiation e.g. to the face back or wrist not being felt in the front of the chest at all.

Character Angina pectoris is classically constricting squeezing pressing or crushing; it is sometimes stinging numbing or burning sometimes it

cannot be described adequately by the patient. It is not sharp shooting or stabbing which are the usual adjectives applied to left inframmary pain. An important characteristic is its constancy the pain being steady while it lasts apart from initial waxing and final waning no pain which is momentary or which repeats itself in a succession of jabs or knife like thrusts is angina.

Duration Attacks are measured in minutes usually they last two or three minutes occasionally five or ten they are not momentary nor do they continue for hours and any pain which behaves in either of these ways is not angina pectoris (as defined above).

Provocation Angina is characteristically produced by any effort that increases the metabolic demands of the myocardium beyond the capacity of the coronary circulation and patients often know or learn the precise amount of effort necessary to provoke pain. When the critical point is reached the patient usually feels compelled to stop whatever he is doing and to stand still until the pain passes off. Attacks are brought on especially by walking uphill or against the wind by hurrying after meals or by any unaccustomed exercise less so by manual work to which the subject is trained. Pain may also be induced by emotion of a kind that raises the blood pressure or increases the cardiac output but this is perhaps less typical emotionally produced pain being more often innocent and associated with anxiety states. In advanced cases pain is provoked by lying down (angina decubitus) or stooping tending to occur when the patient first gets into bed at night or waking him from sleep. It may then depend upon the rise in cardiac output which follows change of posture from vertical to horizontal or upon anxiety dreams.

Pain that occurs after effort but not during it or that is provoked by lying on the left side or by the adoption of some particular posture (other than stooping or lying) is not angina these features are characteristic of left inframmary pain.

DIAGNOSIS OF ANGINA

If a pain conforms in site quality duration and relation to cardiac work to the features mentioned above it is angina pectoris and the diagnosis must stand under any conditions except malingering. The diagnosis should stand likewise when pain conforms to the required features in three out of the four respects mentioned provided it is not untenable in the fourth. For example if a constricting pain brought on only by exertion and lasting but two or three minutes is localised in the left inframmary area it is not angina for the site makes the diagnosis untenable even though it conforms in the other three respects. On the other hand if the same pain is situated between the sternum and the left breast it is almost certainly angina because this site though atypical is not contradictory. Again a midsternal pressing pain brought on only by effort but lasting fifteen minutes is probably angina for the long duration though unusual is not altogether conflicting but should it last two hours it is not angina as defined above.

It is sometimes said that certain associated symptom such as breathlessness, dizziness or faintness, flushing, sweating, weakness and a feeling of impending death help to confirm the diagnosis. It cannot be stressed too strongly that these symptoms carry little weight for they are vasomotor in origin and although they may be provoked by an attack of angina they are in no way characteristic of it and are much commoner in the anxiety states.

The differential diagnosis includes anxiety states, functional disorder or organic disease involving the dorsal spinal ligaments, œsophageal or gastric spasm or distension, diaphragmatic hernia and conditions causing respiratory distress.

Anxiety states with left inframammary pain present no diagnostic difficulty but when pain is parasternal or even central it may be very confusing. The patients are usually women near the menopause and they may describe a central pain radiating to the throat, jaws and arms during or after effort when reaching up to a high shelf when washing or using their arms in other ways and sometimes when emotionally upset. As noted by Cassidy (1946) the attacks are apt to be widely spaced, unrestricted effort causing no distress between them. Complete investigations may reveal nothing significant in any system and the nature of the attacks remains obscure. Angina can only be excluded and then with some uncertainty by obtaining a normal electrocardiogram during spontaneous or induced pain.

Referred pain from the dorsal spinal ligaments may be felt across the front of the chest as in the experimental work of Lewis and Kellgren (1939). Attacks may be related to posture or reproduced by spinal movements or pressure over the interspinous ligaments from D2-4.

Œsophageal spasm may cause central chest pain radiating down both arms and tight or bursting in quality. There is no close relationship to effort and bouts may be periodic like any other gut colic. The diagnosis may be proved by demonstrating œsophageal spasm by means of fluoroscopy and by obtaining a normal electrocardiogram during attack (Wolferth and Edeiken 1942).

Diaphragmatic hernia may cause pain on effort similar to angina pectoris but it is usually more closely related to meals and there may be associated vomiting and belching. Severe attacks may be mistaken for cardiac infarction. The diagnosis is made by means of a barium meal fluoroscopy being carried out with the patient tilted head down (Dwyer 1937).

Relief of pain by belching in any disorder of the œsophagus or stomach is less helpful in distinguishing such conditions from angina pectoris than might be supposed for ischæmic pain may be similarly relieved in about 10 per cent of cases (Riseman and Brown 1937).

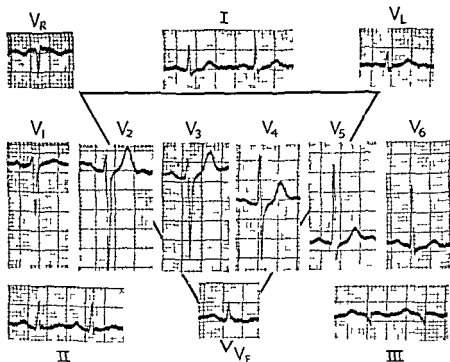
Bronchial asthma or extreme dyspnoea from any cause may be associated with a feeling of substernal tightness that should not be confused with angina pectoris for breathlessness is not a feature of transient myocardial ischæmia.

PHYSICAL EXAMINATION

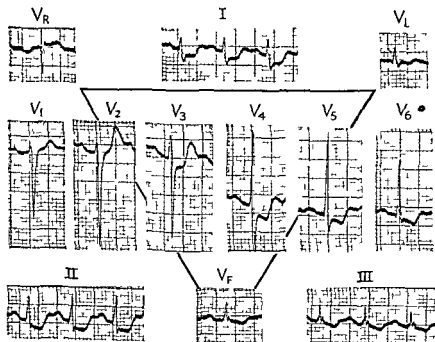
Having made the diagnosis on historical grounds the patient should be examined with a view to ascertaining the cause of the ischæmia. Aortic valve disease and severe anæmia should be recognised by their characteristic features the presence of obesity or of hypertension noted the mental state of the patient assessed and attention should be paid to any other factor which may have any bearing on the frequency or severity of attacks. In this respect diabetes mellitus and polycythæmia must be borne in mind. In the majority of cases however there are no physical signs the rhythm is normal the heart is not enlarged there are no murmurs and there is no evidence of congestive failure the peripheral and fundal arteries and the blood pressure may provide no evidence of general vascular disease fluoroscopy shows a heart shadow normal in size shape and pulsation and the electrocardiogram may be normal at rest. It is repeated for emphasis that this apparent normality of the cardiovascular system is typical of pure angina pectoris due to coronary atherosclerosis and that with few exceptions physical signs radiological changes or electrocardiographic abnormalities are due to complications or associated disease even the demonstration of peripheral atherosclerosis proves little for it is common enough without serious involvement of the coronary vessels and is often missing with advanced coronary disease.

SPECIAL TESTS

Most of the special tests are of little help for the circulation is usually normal at rest. Effort tolerance tests based on the behaviour of the pulse rate and venous pressure are of no value. Reproduction of pain by prescribed effort for purposes of accurate analysis is sometimes useful with a bad witness or pain may be induced to ascertain the prophylactic or curative effect of trinitrin. The only reliable test however is to obtain an electrocardiogram immediately after maximum effort, when characteristic depression of the RS-T segment with or without inversion of the U wave clinches the diagnosis (fig. 14.03). The best method is to make the patient exercise until he is in pain if he stops on account of fatigue or breathlessness without developing pain angina is unlikely especially if the electrocardiogram remains normal. In the author's experience very few (less than 10 per cent) electrocardiograms remain normal during or immediately after an attack of true angina or after sufficient effort to cause breathlessness and fatigue (in ischæmic subjects). The other method is to take serial electrocardiograms while the patient breathes 10 per cent oxygen for twenty minutes or for a shorter time if pain is produced. As depression of the RS-T segment occurs in normal subjects with this test a positive result is only accepted if the depression exceeds 2.5 mm. in any lead or if the T wave becomes inverted in left ventricular surface leads or their counterparts (Levy *et al.* 1938 1939 1941). The test is positive in 3 to 5 per cent of normal controls (Biorck 1946).



(a) ANGINA PECTORIS NO PAIN AT REST



(b) AFTER EFFORT NO PAIN

Fig 14 03—Electrocardiogram (a) before and (b) after exertion sufficient to cause breathlessness and fatigue in a case subject to attacks of angina pectoris: the control record (a) is normal; the second record (b) shows significant depression of the ST segment in practically all leads.

Weintraub and Bishop 1947) in 15 to 20 per cent of cases of doubtful angina, and in 50 to 55 per cent of cases of undisputed angina (Levy *et al* 1941 Biorek, 1946) In the opinion of the writer this test is less useful than the effort test being more difficult to carry out more difficult to interpret more dangerous and far less frequently positive

COURSE

The onset of angina pectoris is more often sudden than gradual and is usually due to a small coronary thrombosis insufficient to cause cardiac infarction The patient may say he was capable of climbing mountains a week ago yet now he can scarcely walk 100 yards Less commonly pain is first experienced during unusually heavy exertion and gradually becomes more easily provoked This represents the slow development of occlusive atherosclerosis

The subsequent course is apt to be punctuated by short periods of relatively sudden deterioration followed by long periods of gradual improvement these episodes signify thrombotic occlusion of a medium sized coronary artery followed by the development of a collateral circulation (Schlesinger 1938) and perhaps by recanalisation

Sooner or later in the majority of cases thrombosis occludes one of the main coronary arteries and cardiac infarction results but a major thrombosis may occur without infarction infarction may occur without thrombosis and ventricular fibrillation may terminate the illness in the absence of both (Appelbaum and Nicolson 1955 Nathanson 1936)

Angina may cause total incapacity in really severe cases and may finally occur at rest (status anginosus or acute coronary insufficiency)

Some cases severe or otherwise improve after cardiac infarction others lose their pain on developing congestive heart failure It is not clear why this should be so but the explanation may be related to the fact that ligation of the coronary vein appears to improve the coronary circulation (Beck and Mako 1941)

PROGNOSIS

The average life expectancy from the onset of angina pectoris is nine to ten years (White Bland and Miskall 1943) about 10 per cent live well might twenty years e.g. John Hunter 1773-93 Sir James Mackenzie 1907-25 Sir Thomas Lewis, 1927-45 Women have a better prognosis than men, and subjects over 40 years of age at the onset fare better than those under 40 (Parker *et al* 1946) Cardiac infarction hypertension enlargement of the heart changes of rhythm bundle branch block and other electrocardiographic abnormalities (at rest) all influence the prognosis adversely (Montgomery, Dry and Gage 1947)

TREATMENT

Conservative The majority of patients with uncomplicated angina of

mild or moderate severity are able to carry out sedentary or light manual work. Any mental or physical activity that increases the frequency of attacks or that causes pain directly should be avoided whilst adequate rest and relaxation should be assured. Diet should be light and its fat content low although hypercholesterolemia is difficult to influence by such means.

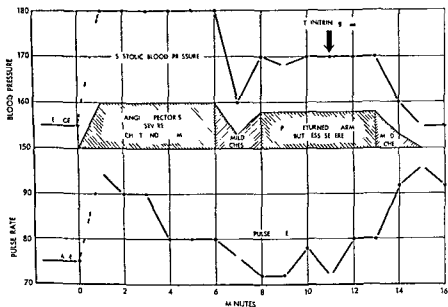


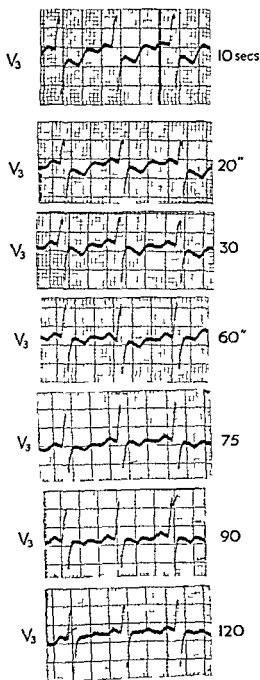
Fig 14.04—Graph showing close correlation between the height of the blood pressure and the degree or extent of pain during an attack of angina pectoris treated with trinitrin.

Smoking should be given up if it is found to contribute to the frequency of attacks. Alcohol in moderation is not harmful in fact as a vasodilator it may be beneficial. Contributory factors such as hypertension, obesity, anaemia, diabetes mellitus, and anxiety should be corrected as far as possible.

Trinitrin 1/100 to 1/120 of a grain (0.5 mg) introduced by Murrell in 1879 may be carried and slipped under the tongue as required either to relieve an attack or before some unavoidable effort which might induce one. Trinitrin is absorbed quickly through the oral mucosa and acts as a coronary vasodilator relieving pain without necessarily altering the blood pressure (Wayne and Laplace 1933-34) but if the blood pressure is lowered as well so much the better (fig 14.04). Ischæmic ST depression in the electrocardiogram is corrected quickly (fig 14.05).

Amyl nitrite 5 minims (0.3 ml) is also effective but less convenient (fig 14.06) a capsule may be broken in a handkerchief and inhaled the noise of the procedure, the pungent smell of the vapour and the

ANGINA PECTORIS



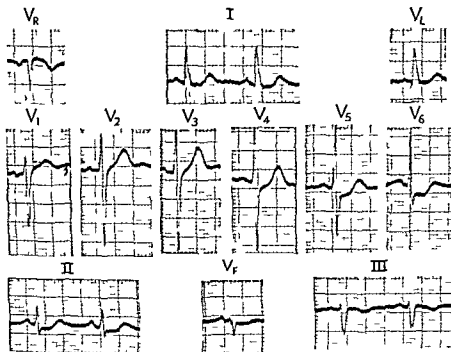
AFTER TRINITRIN

Fig 14 05—Graph illustrating rapid correction of ischaemic depression of the S T segment in lead V₃ in a patient with angina pectoris by means of trinitrin

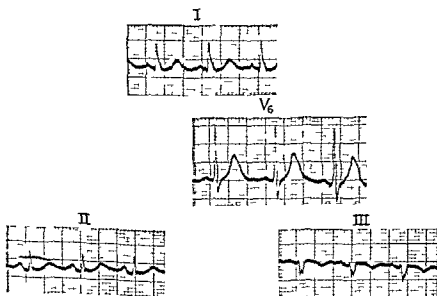
vivid facial flush that accompanies its use, are apt to embarrass the patient in public. Amyl nitrite is a powerful coronary dilator for relief of pain is associated with considerable tachycardia and with conspicuous elevation of the cardiac output. Much interest is also attached to the frequent paradoxical effect of amyl nitrite on the electrocardiogram for the depression of the S T segment that occurs during an attack of angina and which is attributed to subendocardial myocardial ischaemia often becomes further depressed when the drug is inhaled and pain passes off (fig 14 07).

It is doubtful whether any of the drugs used as longer acting coronary vasodilators are of much value (Master Jaffe and Dack 1939). Aminophylline has the best reputation and is employed widely in doses of 0.1 to 0.2 G t d s. It is difficult to demonstrate a physiological effect with such doses but severe angina may be relieved by 0.3 G four hourly if the patient can tolerate it. Epigastric pain and nausea usually prohibit larger doses.

Recent reports have claimed that khellin, an extract from the seeds of an Eastern Mediterranean wild plant *ammi visnaga* is an effective coronary vasodilator with a prolonged action. The dose is 100 mg by mouth three times daily. Angina pectoris is said to be relieved in 74 per cent of cases (Anrep *et al*



AT REST



AFTER AMYL NITRITE

Fig 1406—Electrocardiogram during an attack of myocardial ischaemia treated with amyl nitrite
Expected response showing prompt correction of the depressed S T segment

have been involved in recent years. Few have gained much support but there is something to be said in favour of abolishing pain by means of sensory denervation of the heart achieved by means of section of the upper four dorsal spinal nerve roots or by stellate and upper dorsal ganglion ectomy (White Garvey and Atkins 1933). Destruction of the ganglia by means of alcoholic injection is less certain and may cause intractable root pain in about 10 per cent of cases. Despite the theoretical argument that ganglionectomy may remove nature's warning signal and so allow patients to exercise themselves beyond the limits of safety, there is no doubt that some cases do remarkably well (White and Bland, 1948). Sensory denervation of the heart does not entirely abolish the subjective recognition of an anginal attack although the sensation experienced is not painful. There is good reason to believe also that sympathectomy tends to prevent ventricular fibrillation (Leriche *et al.* 1931; McEachern 1940) and seems to improve the coronary circulation either by preventing reflex spasm (Ivey and Moore 1941) or by causing coronary vasodilatation (Katz and Jochim 1939).

A more drastic surgical procedure aims at improving the coronary circulation by supplying it with a new source of collateral vessels. The idea was based on necropsy observations which showed that the heart might function remarkably well despite almost complete coronary occlusion if for some reason an adequate collateral circulation had developed through the pericardium. These natural results of accident and disease have been marshalled and developed by Claud Beck (1935-36) in the U.S.A. and by O'Shaughnessy (1936-37) in England. Beck sutured a flap of pectoral muscle to the surface of the heart. O'Shaughnessy preferred cardio omentopexy, the omentum being brought up through the diaphragm and stitched or glued on to the surface of the heart after scarification. Whilst experimental evidence affords convincing proof of the establishment of a collateral circulation by such means, the results obtained in clinical cases of ischaemic heart disease scarcely justify the risk entailed.

A simpler means of achieving the same object is to introduce bone dust into the pericardial sac when the pericardial reaction subsides; vascular adhesions offer a collateral source of blood supply to the myocardium (King 1941). This method deserves further trial.

CARDIAC INFARCTION

Myocardial infarction occurs when a mass of heart muscle is sufficiently deprived of its blood supply for an adequate time. The common cause of such an event is coronary thrombosis but coronary embolism, rupture of an atheromatous plaque, subintimal hæmorrhage in an atherosclerotic vessel, dissection and critical lowering of the blood pressure as from shock or hæmorrhage in a patient with occlusive coronary atherosclerosis or syphilitic aortitis may each produce it. Again coronary thrombosis does

not cause myocardial infarction if the collateral circulation is sufficient to preserve the life of the threatened tissue. It follows that coronary thrombosis and myocardial infarction are not synonymous terms and should not be confused: the former means no more than its literal sense implies: the latter means death of a localised mass of heart muscle.

Anatomy of the coronary circulation The site and extent of the infarct depend upon the vessel or vessels occluded, upon the capacity and efficiency of collateral channels, and upon the anatomy of the coronary circulation.

There are two main coronary arteries, left and right. The left divides early into an anterior descending branch and into a left circumflex: the large anterior descending branch runs down the interventricular groove to the apex of the heart and nourishes the anterior part of the right ventricle, the interventricular septum, and the anterior and apical part of the left ventricle; the smaller left circumflex curls round to the back between the left auricle and ventricle and supplies the upper lateral and posterior basal portion of the left ventricle. The right coronary artery does not divide but runs round to the back between the right auricle and ventricle, sending branches to the region of the sinus node, to the anterior part of the right ventricle and to the posterior base of both ventricles. There is a considerable degree of anastomosis between the terminal branches of these vessels, an anastomosis that increases rapidly when the blood supply to any area is threatened (Prinzmetal *et al.* 1947). The right ventricle, supplied as it is by the two biggest coronary arteries and offering little resistance to systolic coronary blood flow, is rarely the seat of infarction. The upper and lateral part of the left ventricle is supplied by proximal branches from both anterior descending and left circumflex vessels and is therefore relatively safe. The posterior basal region is less secure for it is supplied only by terminal branches, some from the right coronary artery and some from the left circumflex. In having this double source of nourishment, however, it is still more fortunate than the anterior apex of the left ventricle, which is fed almost entirely by terminal ramus from the anterior descending branch of the left coronary artery, although anastomotic channels can develop rapidly from the posterior descending branch of the right coronary artery. The interventricular septum is supplied anteriorly by perforating branches from the anterior descending coronary artery and posteriorly by perforating branches from the right. Anastomoses are more conspicuous in the superficial layers of the myocardium than in the inner layers (Prinzmetal *et al.* 1948); they are also at a physiological disadvantage when near the endocardium because they are subjected to a higher intramyocardial pressure (Johnson *et al.* 1939).

Site of thrombosis and infarction Clinically, major coronary thrombosis involves the anterior descending branch of the left coronary artery in 66 to 75 per cent of cases, the right coronary artery in 25 to 40 per cent, and the left circumflex in 5 to 33 per cent (Barnes and Ball 1932; Appelbaum and Nicolson 1935; Munck 1946); thrombosis of the left main trunk is

relatively rare. These figures are conservative for careful study of the whole coronary tree by means of radio opaque injections reveals multiple thromboses in the majority of instances.

The relative incidence of the various sites of infarction harmonises with the anatomical and physiological data and with the sites of thrombosis. In a recent analysis of 160 cases Wartman and Hellerstein (1948) found chiefly anterior infarction in 72 per cent and chiefly posterior in 28 per cent but there were multiple infarcts in 41 per cent. Half the anterior infarcts and a quarter of the posterior infarcts also involved the interventricular septum. Right ventricular infarction rarely occurs alone but it may complicate anteroseptal infarction of the left ventricle.

Combining figures published by Appelbaum and Nicolson (1935) Nathanson (1936) Clawson (1939) and Munck (1946) it is found that coronary thrombosis occurs without cardiac infarction in 20 per cent of cases and that cardiac infarction occurs without coronary thrombosis in 29 per cent in the latter group atherosclerotic occlusion may be complete or incomplete.

Pathology. A cardiac infarct may be difficult to distinguish with the naked eye when less than twenty four hours old microscopically however, acute necrosis of the muscle fibres may be recognised by their swollen appearance and by the loss of their nuclei and striations. When a few days old an infarct is discoloured and may be surrounded by a red zone of hæmorrhage or congestion. Microscopically the necrosed muscle is seen to be invaded by polymorphs. Older infarcts are yellowish white in colour and represent scar tissue.

When necrosis involves the inner layers of the myocardium mural thrombi frequently form against the damaged endocardium in fact they are found in 40 to 50 per cent of all cases (Hellerstein and Martin 1947). Local pericarditis occurs over superficial necrosis and has been reported in 30 to 75 per cent of all cases (Wartman and Hellerstein 1948 Stewart and Turner 1938) diffuse pericarditis develops in about 10 per cent.

Myocardial softening (myomalacia cordis) may result in rupture of the heart (5 to 15 per cent) or in the formation of a cardiac aneurysm (10 to 30 per cent according to published necropsy figures and according to the definition of an aneurysm).

Symptoms. Although the onset of cardiac infarction is sudden premonitory symptoms are common during the preceding week or so and take the form of typical or atypical angina pectoris. Then or without warning of any kind and usually without any obvious precipitating cause the major attack overwhelms the patient. It may strike indiscriminately whether the subject is asleep at rest performing daily routine duties or exerting himself and is commonly signalled by pain indistinguishable in site radiation and quality from angina pectoris but instead of passing off in a few minutes it lasts for hour. Its intensity varies from a feeling of pressure to extreme

agony and gives no indication of the size of the infarct. There may be no other symptoms: on the other hand there may be collapse, weakness, faintness, sweating, pallor, breathlessness and vomiting. Whilst a classical attack is characterised by pain, others present with syncope and yet others with suffocation. In the syncopal type, which represents a vaso-vagal reaction, loss of consciousness may prevent appreciation of pain when paroxysmal cardiac dyspnoea or acute pulmonary oedema dominates the scene: the patient usually admits pain on close questioning.

Physical signs. Unlike angina pectoris, myocardial infarction provides a wealth of physical signs and special findings. When first seen the patient is usually grey, cold, sweating, obviously ill and in pain: he may be breathless and cyanosed, or he may be pale and collapsed—perhaps unconscious. On the other hand he may present none of these features. Within two or three days mild cases may look and feel well.

The jugular venous pressure is sometimes a little raised during the first day or two, and the pulse rate accelerated, but in cases with a vaso-vagal reaction there may be bradycardia. There may be orthopnoea, paroxysmal cardiac dyspnoea or frank pulmonary oedema in severe cases.

The blood pressure falls initially only in cases with a vaso-vagal reaction and indeed may be elevated during the first twelve hours or so (Weiss 1939); in animals it is similarly maintained for the first twenty-four hours (Gross *et al.* 1938) but it drops later, commonly reaching its lowest level on the third or fourth day, when systolic pressures of 80 to 90 mm. of mercury are often found. Thereafter it remains low for several days, or even for weeks, and then in all who survive climbs slowly back towards its previous level, which it may or may not reach (fig. 14.09). In 67 per cent of fatal cases Chambers (1947) observed no such recovery. In hypertensive subjects this drop in pressure may not be recognised unless the original level is known.

The heart sounds are often faint, particularly when the blood pressure is low, and there may be presystolic or protodiastolic gallop rhythm. Transient pericardial friction is heard in about 10 per cent of cases, especially when the infarct is anterior. Disturbances of rhythm are not uncommon and include ectopic beats, paroxysmal ventricular tachycardia, auricular flutter or fibrillation, and any grade of heart block.

Low grade fever is the rule and may continue for several days, but rarely for more than a week. Transient polymorphonuclear leucocytosis also occurs during the first few days. The sedimentation rate begins to accelerate after a day or two, reaches maximum velocity towards the end of the first week, and then gradually returns to normal in an average period of six weeks from the onset (fig. 14.10).

Electrocardiographic appearances. These have already been described and explained in Chapter III (page 93). Leads facing the surface of the infarct show a prominent or monophasic Q wave, initial elevation of the RS-T segment, and subsequent inversion of the T wave. Anterior infarcts may

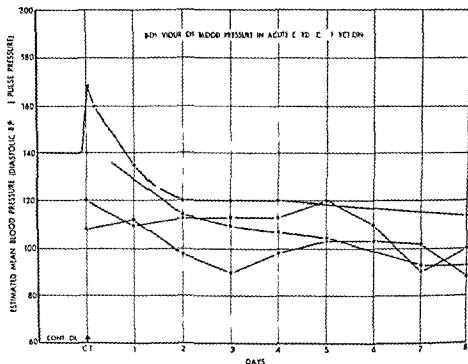


Fig 14 09—Behaviour of the blood pressure in four cases of acute myocardial infarction

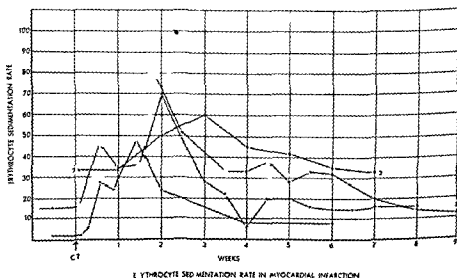


Fig 14 10—Behaviour of the sedimentation rate in four cases of acute myocardial infarction

be mapped out with precision by means of multiple unipolar chest leads and may be chiefly anterolateral (fig 14 11) or anteroseptal (fig 14 12) The Q T pattern is usually transmitted to lead VL and hence mainly to standard lead 1 but if the heart is electrically vertical a V₅ Q T pattern may be transmitted to lead VF, and hence to standard leads 2 and 3 The Q I pattern of posterior infarcts is seen in oesophageal leads over the posterior surface of the left ventricle and is transmitted to lead VF and hence to standard lead 3 (fig 14 13a) while chest leads usually show initial depression of the RS T segment followed by unusually tall T waves (fig 14 13b)

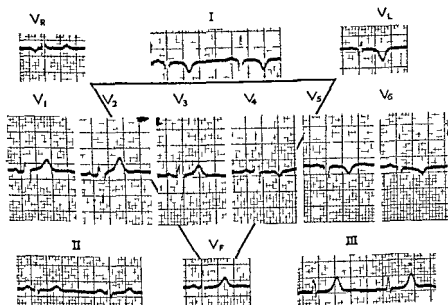


Fig 14 11—Electrocardiogram showing anterolateral cardiac infarction. Maximum changes are seen in leads V₅, V₆, V_L, and standard lead I.

The abnormal Q wave develops early and may persist indefinitely. Elevation of the R T segment is usually transient but a monophasic Q wave associated with persistent elevation of the Q T segment is often seen with ventricular aneurysm (fig 14 14). Primary inversion of the T wave appears in a few days, reaches a maximum within two or three weeks and then gradually reverts towards normal but slight inversion with Pardee coving of the RS T segment may persist in one or more leads (fig 14 15).

The diagnosis of acute cardiac infarction is practically untenable if serial electrocardiograms remain normal in all the recognised leads but an initial electrocardiogram may be normal occasionally if taken within a few hours of the onset.

In differential diagnosis great stress is laid on the abnormal Q wave for

this occurs in no other condition. It must of course be distinguished from a normal Q wave measuring 2 or 3 mm and a monophasic downward deflection in standard lead 3 should not be accepted as a Q wave unless Q is also prominent in standard lead 2 and in lead VF (fig 14 16). Elevation of the RS T segment is also seen in pericarditis (page 342) and opposite large S waves in appropriate leads in left ventricular preponderance (page 426) and left bundle branch block but the contour of the S T segment and the general pattern is different as described elsewhere.

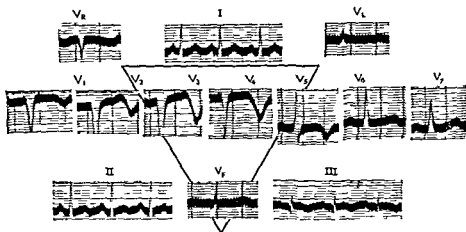


Fig 14 12—Electrocardiogram showing anteroseptal cardiac infarction. Maximum changes are seen in leads V_3 and V_4 .

Primary inversion of the T wave alone is less conclusive evidence of infarction for it may be seen in a variety of conditions including toxic myocarditis, pericarditis, carbon monoxide poisoning, myxœdema, certain biochemical states, and following paroxysmal tachycardia. However, the depth and sharpness of the inversion usually exceed that in all other types, and its association with upward curving of the RS T segment is practically diagnostic. Changes in serial graphs are less helpful because nearly all the primary T wave changes mentioned above are also transient.

Bundle branch block, mostly left, occurred in 7.3 per cent of 700 cases of angina pectoris and in 8.9 per cent of 328 cases of cardiac infarction reported by Salcedo Salgar and White (1935). Conversely, they found that ischaemic heart disease accounted for approximately 50 per cent of 181 cases of intraventricular block of all types. Myster, Dack, and Jaffe (1938) found the incidence of bundle branch block in acute coronary occlusion to be 12 per cent in 1,058 cases collected from the literature and 15 per cent in 375 cases of their own. Intraventricular block does not necessarily imply septal infarction in these cases, and of course may precede the acute episode. Its importance lies in the fact that it may mask the electrocardiographic signs of cardiac infarction, for as explained on page 90, there can be no

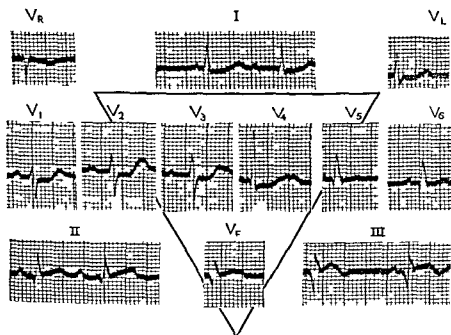


Fig 14.13 (a)—Electrocardiogram showing posterior cardiac infarction. Characteristic changes are seen in leads V_F and hence in leads 2 and 3. The ST segment is depressed in lead V_4 .

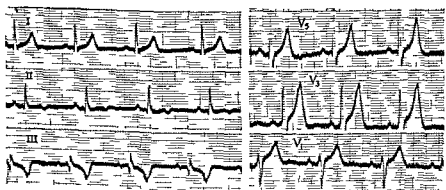
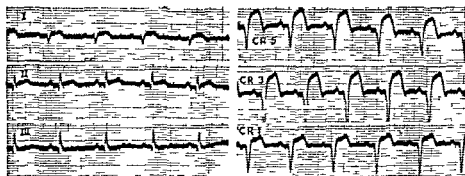
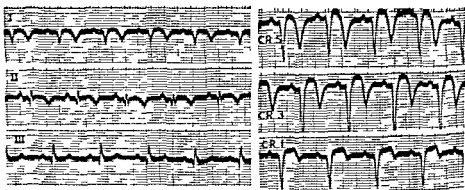


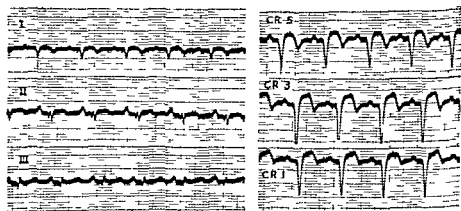
Fig 14.13 (b)—The later stage of posterior infarction showing unusually tall T waves in chest leads.



(a) 29th November 1941



(b) 15th December 1941



(c) 3rd March 1942

Fig 14 14—Electrocardiogram showing widespread monophasic Q waves and persistent elevation of the ST segment associated with ventricular aneurysm

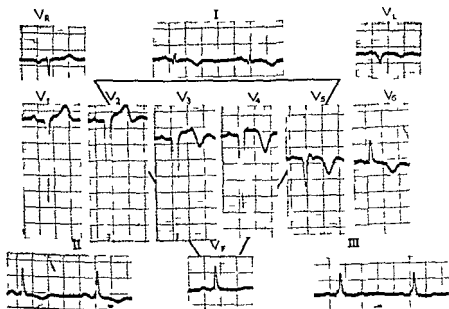


Fig 14 15—Electrocardiogram of a case of old cardiac infarction showing persistent Q waves and Pardee coupling of the ST segment in anterior left ventricular surface leads and their counterparts (leads V_L and standard lead I). The infarct occurred 14 months previously

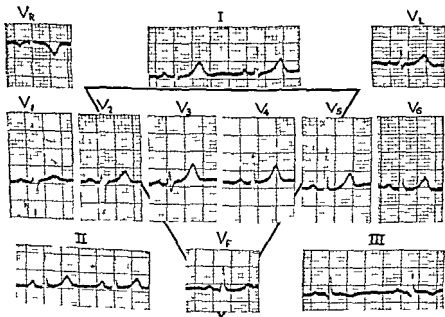


Fig 14 16—Electrocardiogram in a case of pregnancy showing a prominent Q wave and inversion of the T wave in lead 3 due to cardiac rotation, not the absence of a pathological Q wave in lead V_F and the presence of an S wave in standard lead I

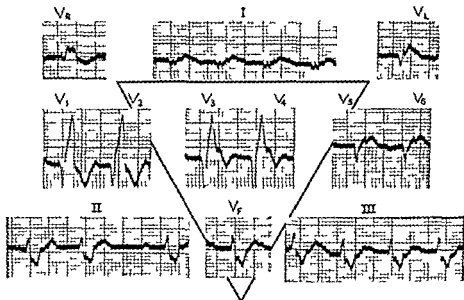


Fig 14 17—Electrocardiogram showing typical appearances of anterior cardiac infarction in the presence of right bundle branch block

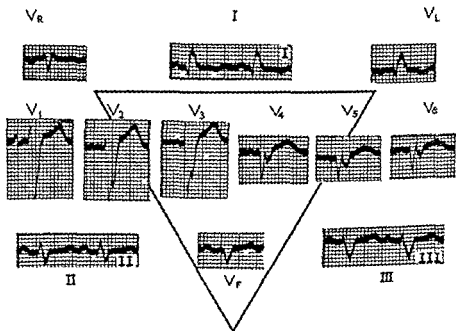


Fig 14 18—Electrocardiogram showing typical appearances of anterior cardiac infarction in the presence of left bundle branch block

Q wave in leads facing the surface of the left ventricle in cases of left bundle branch block unless the septum is also necrosed and the gross deformity of the R-T component may overshadow RS-T changes due to the infarct. Somerville and Wood (1949) however found that the characteristic Q-T pattern of an infarct could be recognised in nearly all cases complicated by right bundle branch block (fig. 14.17) and in about half those with left bundle branch block (fig. 14.18).

Electrocardiography may be of great value in the diagnosis of myocardial infarction months or years after the event: an abnormal Q wave, local dwarfing of R or primary inversion of the T wave in one or more left ventricular surface leads or their counterparts being particularly helpful.

Radiological findings. Fluoroscopy is impracticable during the acute stage of the illness but may be useful later. An infarct on the left border of the heart near the apex may form a ledge (fig. 14.19). In normal hearts pulsation is seen around the whole surface of the left ventricle; in myocardial infarction there may be local absence of pulsation or pulsation may be locally paradoxical: a portion of the ventricle expanding while the rest contracts; this area of absent or paradoxical pulsation represents the infarct and may be seen on the left border of the heart towards the apex or on the diaphragmatic surface of the left ventricle (with the aid of gas in the stomach) for some reason posterior basal infarcts are less easily visualised. Interpretation of pulsation as seen on the fluoroscope is by no means easy and requires considerable experience of normal variation. The kymograph, a simple device for obtaining a permanent skiagraphic record of cardiac pulsation, has been used with some success as an aid in analysing the findings (fig. 14.20); the electrokymograph is even better. But absence of pulsation at the apex may also be seen occasionally in hypertensive heart failure (fig. 14.21).

Ventricular aneurysm is more easily recognised, particularly when situated towards the apex or left lateral border (fig. 14.22). It should not be confused with a dilated left auricle (fig. 14.23) or with an intrapericardial hæmatoma (fig. 14.24). Increased density and unfolding of the aorta due to atheroma, with or without calcification, may be seen in many cases but cannot be regarded as evidence of coronary atherosclerosis: calcified coronary arteries (Snellen and Nauta, 1937) offer more convincing proof but even these do not necessarily signify ischæmic heart disease.

Apart from the changes mentioned, the size and shape of the heart are usually normal in cases of uncomplicated cardiac infarction; enlargement is commonly due to heart failure or to coincident hypertensive heart disease.

Complications. The acute stage lasts on the average for six weeks, during the earlier part of which many complications may arise, the gravest danger being abrupt death from ventricular fibrillation (fig. 14.25); about 10 per cent of all cases die in this way. Other disturbances of rhythm are also relatively common and include ventricular ectopic beats, paroxysmal ven-

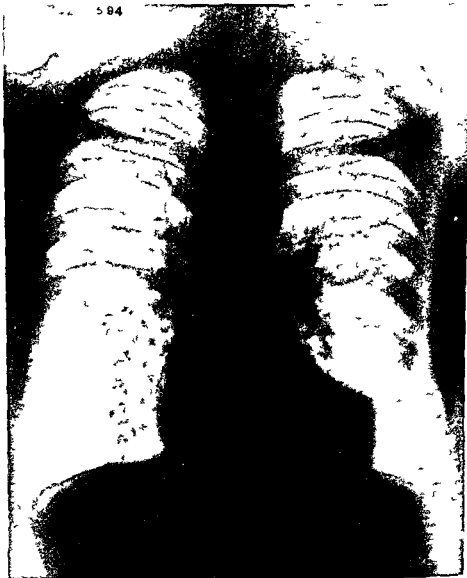


Fig. 14-19—Skiagram in a case of anterior cardiac infarct on showing a ledge on the left border of the heart



Fig 14 20—Kymogram of a case of anterior cardiac infarction showing an area of absent pulsation on the left border of the heart near the apex

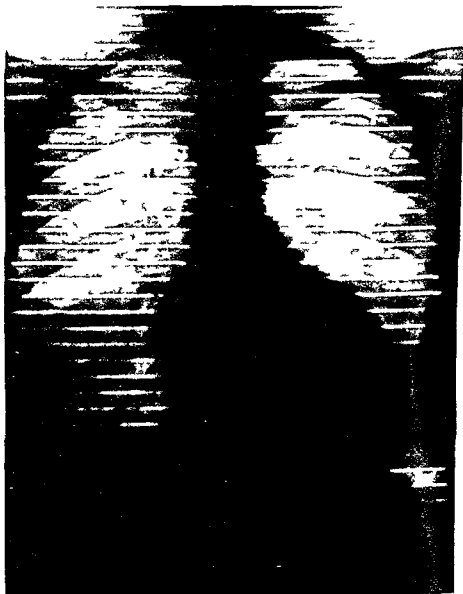


Fig 14-1—Kymogram of a case of hypertensive heart failure showing absence of pulsation at the apex

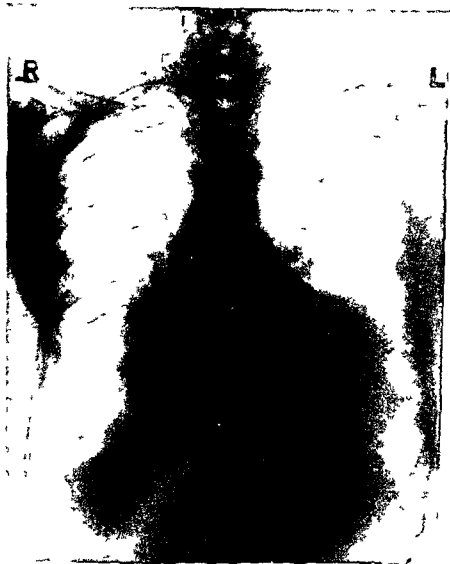


Fig 142.—Skiagram of a case of ventricular aneurysm

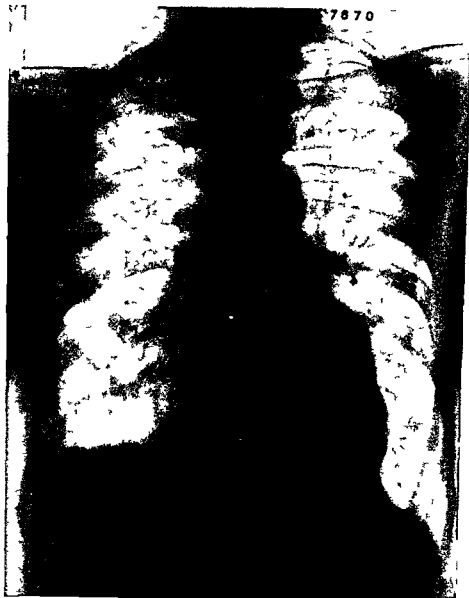


Fig 14 23—Skiagram of a case of organic mitral incompetence showing a dilated left auricle on the left border of the heart

tricular tachycardia paroxysmal auricular flutter and fibrillation nodal rhythm and heart block They should be regarded seriously because they may herald ventricular fibrillation or precipitate heart failure

Left ventricular failure or congestive heart failure is particularly serious and is responsible for as many deaths as ventricular fibrillation more over it increases the risk of phlebothrombosis and pulmonary embolism

Thrombo embolic lesions in various situations are detected clinically in a little over 10 per cent of cases and may be found at necropsy in about 45 per cent (Hellerstein and Martin 1947) The dangerous period is from the fifth or sixth day to the end of the third week when the clotting time is shortened (Ogura *et al* 1946) Phlebothrombosis in the legs resulting in pulmonary embolism is by far the most common and is responsible for



Fig 14 24—Skiagram of a case of stab wound of the heart showing a hæmatoma on its left border (successfully evacuated later)

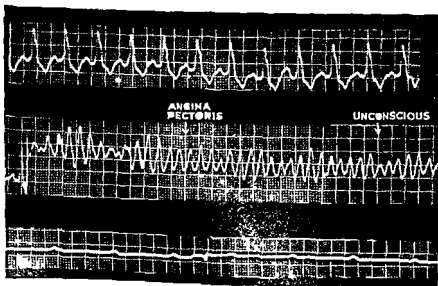


Fig 14 25—Electrocardiogram showing the mode of death in a case of is hæmic heart disease ventricular fibrillation developed while a routine graph was being taken

10 per cent of deaths following cardiac infarction. Cerebral thrombosis or embolism is next in importance accounting for 5 per cent of such deaths. Hellerstein and Martin give the incidence of various thrombo embolic lesions as follows

	<i>Per cent</i>
Pulmonary	23.5
Renal	14.4
Splenic	8.8
Cerebral	7.7
Peripheral arteries	5.5
Mesenteric	1.9
Carotid or aortic	0.5

Although mural thrombi occur in 44 per cent of all cases they are not responsible for more than 10 per cent of these lesions for the latter have been found in 39 per cent of cases without mural thrombi and the total incidence of thrombo embolic lesions is little higher (45 per cent). Thus massive pulmonary embolism is invariably from phlebothrombosis never from mural thrombi and cerebral vascular accidents are commonly the result of local thrombosis (Bean 1938).

Cardiac rupture occurs in less than 1 per cent or in 10 per cent of fatal cases it is not necessarily a dramatic event for the perforation may be small and the signs and symptoms may be those of cardiac compression from hæmopericardium such cases may live a week or more. Perforation of the interventricular septum is seen occasionally and gives rise to the sudden development of a coarse systolic thrill and murmur in the third and fourth intercostal spaces towards the sternum. Heart failure has ensued rapidly in most of the cases reported (e.g. Leonard and Daniels 1938). Although the perforation may look small at necropsy and the track tortuous the shunt during life may be considerable as estimated by means of cardiac catheterisation.

Left ventricular aneurysm may be found at necropsy in as many as 22 per cent of fatal cases (Wartman and Hellerstein 1948) but is less often recognised clinically. In the series referred to above 25 were anterior and 10 posterior five of them ruptured. The condition arises early and may be well developed by the time the patient is allowed up for fluoro copy. The X ray appearances have already been described (page 397). Clinically it is suggested by an unusual pulsation in the region of the apex beat when left ventricular enlargement is improbable on other grounds. The electrocardiogram usually shows a monophasic Q wave and conspicuous and rather persistent elevation of the Q-T segment over the aneurysm while the main QRS deflection is often upright in lead VR (Goldberger and Schwartz 1948) (fig. 14.14). If rupture does not occur during the first few weeks the prognosis is fair.

Pericarditis may be of three kinds (1) a transient friction rub may be heard over an anterior apical infarct and represents local pericardial reaction (2) there may be widespread pericarditis with friction heard at all areas or at a distance from the lesion which may complicate either anterior or posterior infarcts (3) there may be hæmopericardium resulting from ventricular perforation. Local pericarditis does not alter the electrocardiographic pattern of infarction but widespread pericarditis may do so and hæmopericardium invariably does (fig 14 26). Pericardial friction of one kind or another is heard in about 10 per cent of cases.

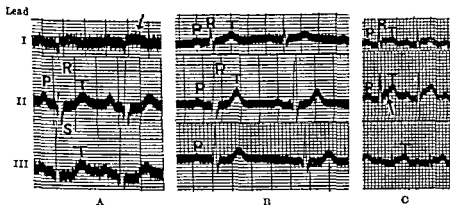


Fig 14 26—Cardiac infarction complicated by perforation and hæmopericardium

A Original anterior cardiac infarction

B After recovery

C After perforation of infarct (hæmopericardium)

After effects The subsequent course is determined by the effect of the occlusion on the total coronary circulation. Angina pectoris may develop or if it was present before it may be worse. On the other hand, if previous pain was due to local ischæmia at the site of the recent infarct, angina may improve or temporarily disappear. Congestive failure may also develop later and may cause the disappearance of angina. Later cardiac rupture is rare and usually denotes fresh coronary occlusion, even when the perforation is through the old infarct. Less than 10 per cent of ruptured hearts are due to an old ventricular aneurysm (Munck 1946).

Differential diagnosis In the differential diagnosis of myocardial infarction many conditions must be borne in mind; the most confusing are massive pulmonary embolism, acute pericarditis, dissecting aneurysm of the aorta, diaphragmatic hernia, œsophageal or gastric dysfunction, and acute pancreatitis, but diaphragmatic pleurisy, especially when bilateral, disease of the gall bladder, perforated duodenal ulcer, epidemic myalgia, and pain referred from the spine may give rise to difficulty. In pulmonary embolism the most important clue is early engorgement of the cervical

veins and immediate hypotension whilst rhythm changes are very rare otherwise both symptoms and signs may be indistinguishable from those of coronary thrombosis and even limb lead electrocardiograms may resemble those of posterior cardiac infarction. Fortunately however chest lead graphs are diagnostic (page 450). Acute pericarditis may simulate cardiac infarction closely but may be distinguished by the electrocardiogram (page 342). Dissecting aneurysm is characterised by radiation of pain to the back and downwards by hypertension by the absence of electrocardiographic change by the development of aortic incompetence and perhaps by signs of involvement of carotid subclavian renal or femoral arteries (page 523). Diaphragmatic hernia should be considered when there are no changes in temperature white count ESR and electrocardiogram and may be diagnosed by means of a barium meal with the patient in the head down position. Oesophageal or gastric pain may be felt in the centre of the chest and may resemble the pain of cardiac infarction but physical examination is entirely negative the electrocardiogram remains normal and the subsequent course is benign. Acute pancreatitis may be recognised by the urinary diastase test.

Treatment Patients should be confined to bed at once and should remain there for three to six weeks or longer according to the severity of the illness and to the behaviour of the sedimentation rate and electrocardiogram. If the blood pressure is low and the patient faint or dizzy he may have to lie flat otherwise and particularly if there is any sign of failure he should be propped up against a back rest in order to reduce the work of the heart (page 158).

Semi starvation for the first few days followed by an 800 calorie diet during the dangerous period practically halves the mortality rate (Master *et al* 1936). Fruit drinks and soft stewed or fresh fruit with sugar is all that should be allowed for the first forty eight hours. The quality of the later light diet matters less than its bulk and calorific value but should contain little sodium if there is any evidence of failure.

The most beneficial drug in the acute phase is morphine which should be given in adequate doses and as often as required to relieve pain and distress and to induce rest and sleep. Excellent results are obtained when pain is severe by giving it intravenously in a dose not exceeding $\frac{1}{4}$ of a grain (15 mg) dissolved in at least 2 ml of sterile water or saline and at a slow rate three minutes being taken over the injection.

Quinidine 3 to 5 grains (0.25 G) t d s may be given in the hope of preventing ventricular fibrillation and other changes of rhythm. Results are difficult to assess but on the whole seem to be encouraging certainly quinidine prevents ventricular fibrillation in dogs (Wegria and Nickerson 1943) and the doses recommended are without danger.

Heparin and dicoumarol (page 454) have been used widely in recent years to prevent extension of coronary thrombosis and thrombosis elsewhere. The frequency of pulmonary embolism and of cerebral thrombosis

has already been noted the former was directly responsible for 6.5 per cent of deaths in a series of 200 fatal cases reported by Eppinger and Kennedy (1938) was present in 24.5 per cent of the total and in 32.7 per cent of those with congestive failure. These are average figures and provide good grounds for anticoagulant therapy. The results of a preliminary analysis of 500 cases so treated (collected by a special committee of the American Heart Association) have been given by Vander Veer, Marshall and Luo (1948) as follows:

	<i>Controls per cent</i>	<i>Cases treated with anticoagulants per cent</i>
Death rate	23	13
Thrombo embolism	19	9

The coronary vasodilators, with the possible exception of aminophylline, do not relieve the pain of cardiac infarction and do not influence its course. aminophylline 0.2 G t.d.s. or four hourly may perhaps improve the collateral circulation and may help to prevent cardiac asthma.

The only other drugs used at all frequently are bromide and phenobarbitone to allay anxiety in apprehensive patients.

Adrenaline should be avoided, no matter how low the blood pressure, for the gravest danger is ventricular fibrillation, and the drug most likely to produce it. Digitalis and strophanthin are rarely indicated and are dangerous for the same reason (Travell, Gold and Modell, 1938). They may be considered, however, when congestive heart failure is becoming serious or when auricular fibrillation or flutter with rapid ventricular rate persist for more than twenty-four hours, but especial care must be taken to avoid an overdose. Coramine, strychnine, cardiazol and other similar remedies of the stimulant class are not advised.

Other measures may be necessary when there are complications: mercurial diuretics and a low sodium diet are useful in the event of failure; ephedrine $\frac{1}{2}$ gr. (32 mg.) t.d.s. should not be withheld if Stokes-Adams attacks complicate heart block and bolder doses of quinidine may be needed to combat paroxysmal ventricular tachycardia.

If the course is benign and the patient looks and feels well, he may be allowed up after three weeks, provided the sedimentation rate has returned to normal and the electrocardiogram does not show a large infarct. Most cases require a month in bed and a further fortnight resting at home on a couch, but those with complications should remain in bed for six weeks or longer.

Six weeks to three months convalescence is usually needed while the patient regains his confidence and gradually resumes his ordinary activities. Radical change of employment is rarely practicable in this age group, but lighter work and less responsibility may have to be advised. Relatively good

recovery from the first attack is the rule but severe angina or recurrent congestive failure may cause total incapacity after subsequent attacks

Prognosis The mortality rate in acute cardiac infarction has been about 25 per cent for all attacks not specially treated with prompt nursing attention initial semi starvation and no interference it is said to be 16.5 per cent (Master *et al* 1936) and with anticoagulant therapy 13 per cent (Vander Veer *et al* 1948) With the added help of a low sodium diet and quinsidine the mortality rate for all attacks should not exceed 10 per cent Such figures exclude cases of sudden death due to coronary thrombosis they refer only to those which survive long enough to receive medical attention The mortality rate in first attacks has been about two thirds that in all attacks

Preceding angina pectoris or hypertension has no influence on the prognosis but diabetes mellitus is adverse Whether the infarct is anterior or posterior matters little but involvement of the septum multiple lesions and extensive necrosis as judged by the electrocardiogram are bad omens Initial shock prolonged hypotension a small pulse pressure arrhythmias cardiac enlargement congestive heart failure and thrombo embolism are naturally serious (Katz and Mintz 1947) Both in hypertensive and normotensive subjects the greater the fall in blood pressure the worse the outlook again those whose blood pressure fails to climb back towards their previous levels fare badly (Chambers 1947)

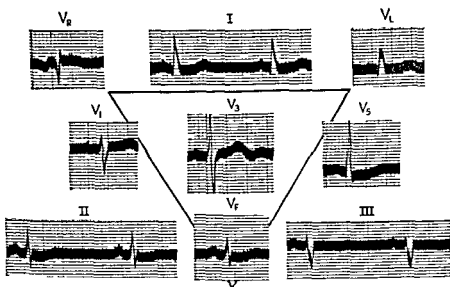
The further outlook in those who make a satisfactory recovery from their first attack of cardiac infarction does not differ radically from that in angina pectoris as a whole reliable figures are not available but the average life expectancy may be estimated at about seven to eight years

The factors which influence the ultimate prognosis are also the same as those in angina (page 380)

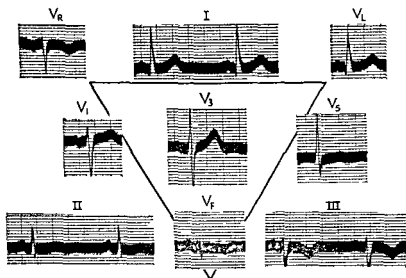
ACUTE CORONARY INSUFFICIENCY

The term acute coronary insufficiency has been proposed by Master and his colleagues (1947) to describe those cases of ischaemic heart disease that cannot properly be called angina pectoris or cardiac infarction but that appear to be something between the two

The physiological basis for the condition is similar to that for angina pectoris but the responsible precipitating factors are prolonged For example prolonged increase of cardiac work may be due to paroxysmal tachycardia auricular flutter hypertensive crises thyrotoxic crises and to an overdose of certain drugs such as adrenaline prolonged diminution of the coronary blood flow may be due to anything that seriously lowers the cardiac output and blood pressure e.g. hæmorrhage shock massive pulmonary embolism and vaso-vagal syncope prolonged reduction of the oxygen content of the arterial blood results from asphyxia carbon monoxide poisoning and acute anæmia In all these conditions the nutritional de-



AT REST WHEN NOT IN PAIN



AFTER REST AND DICOUMAROL

Fig. 14-27—Electrocardiogram showing persistent depression of the ST segment in a case of acute coronary insufficiency. Recovery followed 4 weeks rest and dicoumarol.

mands of the myocardium may be inadequately met especially if occlusive coronary atherosclerosis is also present and subendocardial necrosis of any part of the left ventricle may occur. The inner third of the muscle suffers most as in angina pectoris owing to the intramyocardial pressure gradient.

The clinical features may resemble a prolonged attack of angina or atypical cardiac infarction; on the other hand the condition may be clinically silent. The electrocardiogram shows depression of the RS-T segment in left ventricular surface leads and their counterparts similar to that seen during an attack of angina pectoris or during artificial hypoxia (fig. 14 27)

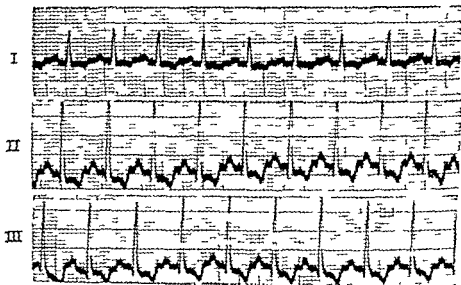
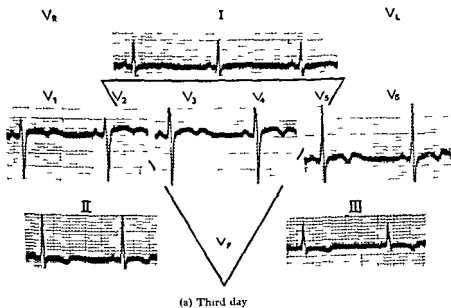


Fig. 14 28—Electrocardiogram showing transient inversion of the T waves following prolonged circulatory collapse with extreme tachycardia without evidence of structural disease of the heart.

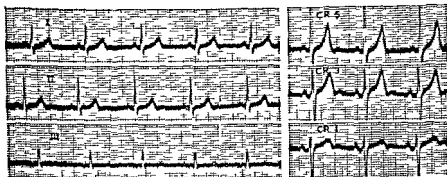
Transient inversion of the I wave in the same leads may develop later in the attack and may last for several days after the circulation has returned to normal (fig. 14 28). True inversion of the I waves is well seen in carbon monoxide poisoning (fig. 14 29).

Acute coronary insufficiency as so defined may explain the occurrence of true heart failure in the later stages of shock (Wiggers 1947) and the readiness with which heart failure develops during infusions in the hyperkinetic stage of severe haemorrhage or acute anaemia (Sharpey-Schafer 1944). It may also explain the poor functional state of the heart too long subjected to tamponade and the development of true heart failure in occasional cases of constrictive pericarditis.

A special type of acute coronary insufficiency occurs after coronary thrombosis without cardiac infarction. This is described on page 84.



(a) Third day



(b) Twelve days later

Fig 14-29—Transient inversion of the T waves due to carbon monoxide poisoning.

REFERENCES

- Anrep G V Barsoum G S Kenawy M R and Misrahy G (1946) Ammi vi naga in the treatment of the anginal syndrome *Brit Heart J* 8 171
 ——— (1947) Therapeutic uses of khellin Method of standardisation *Lancet* i 557
- Appelbaum E and Nicolson G H B (1935) Occlusive diseases of coronary arteries analysis of pathological anatomy of 168 cases with electrocardiographic correlation in 36 of these *Amer Heart J* 10 662
- Barnes A R and Ball R G (1932) The incidence and situation of myocardial infarction in one thousand consecutive post mortem examinations *Amer J med Sc* 183 215
- Bean W B (1938) Infarction of the heart 3 Clinical course and morphological findings *Ann intern Med* 12 71
- Beck C S (1935) The development of a new blood supply to the heart by operation *Ann Surg* 102 901 — (1936) Further data on the establishment of a new blood supply to the heart by operation *J thoracic Surg* 5 604 — and Mako A F (1941) Venous stasis in the coronary circulation *Amer Heart J* 21 767
- Ben Asher S (1947) Further observations on the treatment of the anginal syndrome with thiouracil *Ibid* 33 490
- Biorck G (1946) Anoxæmia and exercise tests in the diagnosis of coronary disease *Ibid* 32 689 — (1946) Hypoxæmia tests in coronary disease *Brit Heart J* 8 17
- Blumgart H L Levine S A and Berlin D D (1933) Congestive heart failure and angina pectoris the therapeutic effect of thyroidectomy on patients without clinical or pathological evidence of thyroid toxicity *Arch intern Med* 51 866
- Cassidy M (1946) Coronary disease *Lancet* ii 567
- Chambers W N (1947) Blood pressure studies in 100 cases of coronary occlusion with myocardial infarction *Amer J med Sc* 213 40
- Clawson B J (1939) Coronary sclerosis An analysis of nine hundred and twenty eight cases *Amer Heart J* 17 387
- Cowdry E V (1933) Arteriosclerosis A survey of the problem New York
- Cutler F C and Schnitzer M T (1934) Total thyroidectomy for angina pectoris *Ann Surg* 100 578
- Duguid M B (1946) Thrombosis a factor in the pathogenesis of coronary atherosclerosis *J Path and Bact* 58 207
- Dwyer M F (1937) Hernia of cardiac end of stomach through diaphragm *Radiology* 28 315
- Eppinger E C and Kennedy J A (1938) The cause of death in coronary thrombosis with special reference to pulmonary embolism *Amer J med S* 195 104 — Levine S A (1934) The effect of total thyroidectomy on the response to adrenaline *Proc Soc exper Biol and Med* 31 485
- Goldberger E and Schwartz S P (1948) Electrocardiographic patterns of ventricular aneurysm *Amer J med* 4 243
- Gordon H (1947) Mechanism of lipophage deposition in atherosclerosis *Arch Path* 44 247
- Gordon W H Bland E F and White P D (1939) Coronary artery disease analysed post mortem *Amer Heart J* 17 10
- Gross L Schauer G and Mendlowitz M (1938) Hemodynamic studies in experimental coronary occlusion *Ibid* 16 278
- Harrison C V and Wood P H (1949) Hypertensive and ischemic heart disease a comparative clinical and pathological study *Brit Heart J* 11 205
- Heberden W (1801) Commentaries on the history and cure of diseases London

- Hedley O F (1939) Analysis of 5 116 deaths reported as due to acute coronary occlusion in Philadelphia 1933-7 *Public Health Rep* Washington p 972
- Hellerstein H K and Martin J W (1947) Incidence of thromboembolic lesions accompanying myocardial infarction *Amer Heart J* 33 443
- Herrick J B (1912) Clinical features of sudden obstruction of the coronary arteries *J Amer med Ass* 59 2015
- Hunter J (1796) A treatise on the blood inflammation and gunshot wounds Philadelphia
- Johnson J R., and Di Palma J A (1939) Intra myocardial pressure and its relations to aortic blood pressure *Amer J Physiol* 125 234
- Katz L N and Jochim K (1939) Observations on innervation of coronary vessels of dog *Ibid* 126 395
- Mintz S S (1947) An analysis of immediate mortality in 572 cases of recent myocardial infarction *J lab clin Med* 32 325
- King E S J (1941) Surgery of the heart Baltimore
- Leary T (1934) Experimental atherosclerosis in the rabbit compared with human (coronary) atherosclerosis *Arch Path* 17 453 — (1938) Vascularisation of atherosclerotic lesions *Amer Heart J* 16 549
- Lehr D (1948) Lowered incidence of sensitisation through the use of sulpho namide combinations *Brit med J* ii 4576
- Leonard B W and Daniels W B (1938) Perforation of the interventricular septum caused by coronary occlusion *Amer Heart J* 16 751
- Leriche R Hermann L and Fontane R (1931) Ligature de la coronaire gauche et fonction cardiaque chez l'animal intact *CR Soc Biol* 107 545
- Levy R L Barach A L and Bruenn H G (1938) Effects of induced oxygen want in patients with cardiac pain *Amer Heart J* 15 187 — Bruenn H G and Russell N G (1939) The use of the electrocardiographic changes caused by induced anoxæmia as a test for coronary insufficiency *Amer J med Sc* 197 241 — and Moore R L (1941) Paravertebral sympathetic block with alcohol for relief of cardiac pain report of 45 cases *J Amer med Ass* 116 2563 — Williams N E Bruenn H G and Carr H A (1941) The anoxæmia test in the diagnosis of coronary insufficiency *Amer Heart J* 21 634
- Lewis T (1934) Clinical science London — Kellgren J H (1939) Observations relating to referred pain visceromotor reflexes and other associated phenomena *Clin Sc* 4 47
- McEachern C G Manning G W and Hall G E (1940) Sudden occlusion of coronary arteries following removal of cardio sensory pathways an experimental study *Arch intern Med* 65 661
- McNee J W (19-5) Clinical syndrome of thrombosis of coronary arteries *Quart J Med* 19 44
- Master A M (1936) Treatment and immediate prognosis of coronary artery thrombosis 267 attacks *Amer Heart J* 12 549 — (1947) Incidence of acute coronary artery occlusion *Ibid* 33 135 — Grisham A Field L E and Horn H (1947) Acute coronary insufficiency an entity *J Mount Sinai Hosp* 14 8 — Dack S and Jaffe H L (1938) Bundle branch and intraventricular block in acute coronary artery occlusion *Amer Heart J* 16 783 — Jaffe H L and Dack S (1939) The drug treatment of angina pectoris due to coronary artery disease *Amer J med Sc* 197 774
- Montgomery G E Dry Th J and Gage R P (1947) Further observations on the prognosis in angina pectoris due to coronary sclerosis *Minnesota Med* 30 162
- Munck W (1946) Pathological anatomy of sudden heart death *Acta Path et Micro Scand* 23 107
- Murrell W (1879) Nitro glycerine as a remedy for angina pectoris *Lancet* i 80

Nathanson M H (1936) Pathology and pharmacology of cardiac syncope and sudden death *Arch intern Med* 58 685

Newman M (1946) Coronary occlusion in young adults. A review of fifty cases in the Services *Lancet* ii 409

Opura J H Fetter N R Blankenhorn M A and Glueck H I (1946) Changes in blood coagulation following coronary thrombosis measured by the heparin retarded clotting test (Vaugh and Ruddick Test) *J clin Invest* 25 586

O'Shaughnessy I (1936) An experimental method of providing a collateral circulation to the heart *Brit J Surg* 23 663 — (1937) Surgical treatment of cardiac ischaemia *Lancet* i 185 — Slome D Surgical revascularisation of the heart *Ibid* i 617

Parker R L Dry T J Willius F A and Gage R P (1946) Life expectancy in angina pectoris *J Amer med Ass* 131 93

Parry C H (1799) An inquiry into the symptoms and causes of the syncope anginosa commonly called angina pectoris illustrated by dissections London

Paterson J C (1936) Vascularisation and haemorrhage of intima of arterio sclerotic coronary arteries *Arch Path* 22 313 — (1939) Capillary rupture with intimal haemorrhage a cause of pulmonary thrombosis *Amer Heart J* 18 451 — (1941) Some factors in causation of intimal haemorrhages and in precipitation of coronary thrombi *Canad med Ass J* 44 114

Poe W D (1947) Fatal coronary artery disease in young men *Amer Heart J* 33 76

Prinzmetal M Bergman H C Kruger H E Schwartz L L Simkin B and Sobin S S (1948) Studies on the coronary circulation III Collateral circulation of beating human and dog hearts with coronary occlusion *Ibid* 35 689 — Simkin B Bergman H C and Kruger H E (1947) Studies on the coronary circulation II The collateral circulation of the normal human heart by coronary perfusion with radioactive erythrocytes and glass spheres *Ibid* 33 4-0

Raab W (1945) Thiouracil treatment of angina pectoris *J Amer med Ass* 128 439

Riseman J E F and Brown M G (1937) Analysis of diagnostic criteria of angina pectoris critical study of 100 proved cases *Amer Heart J* 14 331

Ryle J A and Russell W T (1949) The natural history of coronary disease A clinical and epidemiological study *Brit Heart J* 11 370

Salcedo Salgar J and White P D (1935) Relationship of heart block auriculo ventricular and intraventricular to clinical manifestations of coronary disease angina pectoris and coronary thrombosis *Ibid* 10 1067

Sharpey Schafer E P (1944) Circulatory dynamics of haemorrhage *Brit med Bull* 2 171

Schlesinger M J (1938) An injection plus dissection study of coronary artery occlusions and anastomoses *Amer Heart J* 15 5-8

Shute E V (1945) Effect of vitamin F upon impaired kidney function (preliminary note) *Canad med Ass J* 52

Snellen H A and Nauta J H (1937) Roentgen diagnosis of coronary calcification *Fortschr a d Geb d Roentgenstrahlen* 56 277

Somerville W and Wood P H (1949) The electrocardiogram of cardiac infarction associated with bundle branch block *Brit Heart J* 11 305

Stewart C F and Turner L B (1938) A note on pericardial involvement in coronary thrombosis *Amer Heart J* 15 23-

Travell J Gold H and Modell W (1938) Effect of experimental cardiac infarction on response to digitals *Arch intern Med* 61 154

Van der Veer J B Marshall D S and Kuo P T (1948) Experiences with the use of heparin and dicoumarol in the treatment of coronary thrombosis and thrombo embolic disease *Trans Coll Phys Philadelphia* 16 67

Wartman W B (1938) Occlusion of the coronary arteries by hæmorrhage into their walls *Amer Heart J* 15 459 — Hellerstein H K (1948) The incidence of heart disease in 2 000 consecutive autopsies *Ann intern Med* 28 41

Wayne E J and Laplace I B (1933-4) Observations on angina of effort *Clin Sc* 1 103

Wegria R and Nickerson N D (1943) The benzol adrenaline test as a reliable method of estimating changes in the sensitivity of the dog's ventricles to fibrillation. Application of the method to the study of quinidine sulfate *Amer Heart J* 25 58

Weintraub H J and Bishop L F (1947) The anoxæmia test for coronary insufficiency *Ann intern Med* 26 741

Weiss M M (1939) The early rise of blood pressure in coronary thrombosis *Amer Heart J* 17 103

White J C and Bland E F (1948) The surgical relief of severe angina pectoris. Methods employed and end results in 83 patients *Medicine* 27 1 —

Garrey W E and Atkins J A (1933) Cardiac innervation: experimental and clinical studies *Arch Surg* 26 765

White P D, Bland E F and Miskall E W (1943) The prognosis of angina pectoris: a long time follow up of 497 cases including a note on 75 additional cases of angina pectoris decubitus *J Amer med Ass* 123 801

Wiggers C J (1947) Myocardial depression in shock *Amer Heart J* 33 633

Wilens S L (1947) The relationship of chronic alcoholism to atherosclerosis *J Amer med Ass* 135 1136

Wolferth C C and Edeiken J (1942) The differential diagnosis of angina pectoris with special reference to œsophageal spasm and coronary occlusion *Pennsylvania med J* 45 579

CHAPTER XX

HYPERTENSIVE HEART DISEASE

HYPERTENSIVE heart disease is but one facet of the whole problem of systemic hypertension. It is necessary to consider this problem first.

DEFINITION

Hypertension implies elevation of the basal blood pressure above the normal limits of 145/90 mm Hg. Physiological vasoconstriction due to emotion, to cold, or to other trivial cause is common and modifies the significance of casual high readings of the order of 160/90 mm Hg. The basal pressure is that obtained when the subject is lying down and when successive readings at five minute intervals have dropped to a steady level. Strictly speaking, the subject should have had nothing to eat or drink for twelve hours and the room temperature should be about 70° F, but these points are impracticable.

When elevation of the blood pressure shows disproportion between systolic and diastolic levels, systolic bias favours rigidity of the aorta and large vessels as in atherosclerosis, or increased force of cardiac contraction as in thyrotoxicosis, whereas diastolic bias favours vasoconstriction as in true hypertension. It is therefore permissible to speak of systolic or diastolic hypertension.

VARIETIES OF HYPERTENSION

Hypertension may be paroxysmal as in pheochromocytoma of the adrenal medulla, transient as in acute nephritis and toxæmia of pregnancy, or persistent as in chronic nephritis, chronic pyelonephritis, surgical kidney, coarctation of the aorta, and essential and malignant hypertension. High blood pressure accompanying thyrotoxicosis and the climacteric is coincidental; statistical analysis shows no significant correlation and the pressure does not fall when these disorders are corrected (Bechgaard 1946). The blood pressure in obese subjects may appear to be higher than it really is, owing to the unreliability of the cuff method of measurement when applied to a fat limb; lower pressures may be recorded by direct arterial puncture. Under certain conditions, e.g. during a rigor when there is intense vasoconstriction, or when the main artery to the limb is partly occluded, the blood pressure reading may be much lower when measured by the cuff method than when measured by direct arterial puncture; indeed it may be immeasurable by ordinary means when direct puncture proves it to be in the region of 100 mm Hg. Such fallacies must be constantly

borne in mind Hypertension associated with mitral stenosis is almost certainly a matter of chance, apart from the transient rise of pressure which may result from heart failure

INCIDENCE

In 1928 hypertension accounted for 14.8 per cent (Bell and Clawson) to 20 per cent (Fahr) of all deaths in the U S A in people over 50 years of age British estimates are similar About 5 per cent of young adults 30 to 40 per cent of subjects over 40 and 65 to 75 per cent of those over 70 have some degree of hypertension (Master *et al* 1943) the lower figures apply to men the higher to women At least 80 per cent of hypertensive subjects are between 40 and 70 years of age the peak period being 50 to 59 (Janeway 1913 Bechgaard 1946) According to Platt (1948) severe persistent hypertension in persons under 40 years of age is commonly renal less than a third of his series were essential and he encountered no primary malignant cases under the age of 34

The sex incidence is about equal men being rather more frequently affected in the upper classes (Janeway 1913 Ehrstrom 1918) women in the lower (Blackford *et al* 1930 Bechgaard 1946) Malignant hypertension however affects three men to one woman

About 80 to 85 per cent of cases of persistent hypertension are essential about 2 per cent are primary malignant and most of the remainder are renal

High blood pressure appears to be linked with civilisation it is said to be rare or uncommon in China amongst orientals generally (Harris 1927) and in negroes (Donnison 1929) but it is common or more common in civilised negroes in the U S A as in the white population (Fishberg 1939)

PATHOGENESIS

Paroxysmal hypertension is due to an excess of circulating adrenaline released by a pheochromocytoma of the adrenal medulla (Beer King and Prinzmetal 1937)

Transient hypertension in acute nephritis appears to depend upon a nervous rather than a humoral agent (Pickering 1943) and may be due to extra renal factors (Kylin 1926) There is reason to believe that acute nephritis is an allergic vascular reaction to the products of remote bacterial infection (Cavelti and Cavelti 1945) usually but not exclusively streptococcal the brunt of the attack falling on the glomerular tufts but the capillaries elsewhere not escaping entirely General vasospasm may cause the hypertension

Hypertension in *toxæmia of pregnancy* may be transient and behave like that in acute nephritis or it may be persistent and resemble essential or malignant hypertension (Golden Dexter and Weiss 1943)

High blood pressure in *coarctation of the aorta* (page 208) probably results from diminution of the renal blood flow. It does not occur experimentally if the aorta is constricted below the origin of the renal arteries (Rytand 1938).

Hypertension resulting from *chronic nephritis*, *chronic pyelonephritis* (Schoen 1930 Longcope and Winkenwerder 1933) and certain *surgical kidneys* (Braasch Walters and Hammer 1940) is almost certainly attributable to a humoral agent liberated by the diseased kidney (Pickering 1943).

ETIOLOGY OF ESSENTIAL HYPERTENSION

Certain predisposing factors must be considered first.

Heredity According to Platt (1947) essential hypertension could be a hereditary disease conveyed as a Mendelian dominant with a rate of expression of more than 90 per cent. This may be an extreme view but the importance of the hereditary factor cannot be denied. Thus Ayman (1934) studying 277 families found hypertension in the children in 3.1 per cent of the families when both parents were normal, in 28.3 per cent when one parent was hypertensive and in 45.5 per cent when both parents were hypertensive. Again in an investigation based upon 236 members of 30 families Hines (1940) found that the children were hyper reactors to the cold pressor test in 43.4 per cent when one parent was either hypertensive or a hyper reactor and in 95 per cent when both parents were affected. In Bechgaard's series of over 1000 cases of persistent hypertension which included 20.7 per cent possible renal cases (in which there is no hereditary factor) one or both parents were seriously hypertensive in 75 per cent.

Hyper reaction to pressor agents The excessive reaction of hypertensives to the cold pressor test of Hines (1940) is the best example. The test is carried out as follows: the basal blood pressure is first recorded in the usual way; the subject's free hand is then plunged into ice cold water (3 to 5°C) to just above the level of the wrist and immersed for one minute while the blood pressure is recorded at half minute intervals. In 85 per cent of normal persons the blood pressure rises an average of 12.4/10.1 mm Hg and returns to its previous level within two minutes. A rise of more than 20/15 mm Hg is regarded as a hyper reaction. Patients with established essential hypertension show an average rise of 46.6/30.9 mm Hg, 95 per cent being hyper reactors. Follow up studies indicate that apparently normal individuals who are hyper sensitive to the cold pressor test are likely to develop persistent hypertension. Hines also claims that high casual readings due to emotion have the same significance but this is not substantiated by the subsequent histories of patients with Da Costa's syndrome (Grant 1925 Wood 1941).

Holding the breath for 20 seconds may also be used as a pressor agent.

in much the same way, and compares favourably with the cold pressor test (Ayman and Goldshine 1939)

Other factors The influence of civilisation and of sex in malignant hypertension has already been mentioned

Structural changes in the vessels Certain structural vascular changes often found associated with hypertension have been proved to play no part in its production. Atherosclerosis is innocent in this respect unless a plaque constricts the renal artery, increased rigidity of the aorta and great vessels may raise the systolic pressure, increase the pulse volume and accelerate the speed of the pulse wave but it has little influence upon the mean blood pressure. Calcification of the media of medium sized arteries has a similar effect. The characteristic vascular lesion which is the signature of malignant hypertension, necrosing afferent glomerular arteriolitis, is a result, not a cause of extreme hypertension. Multiplication of the internal elastic lamina and hypertrophy of the media of small arteries and arterioles are also effects, not a cause of sustained hypertension. Hyaline thickening of the intima, especially of the afferent glomerular arteriole, found in 98 per cent of cases of essential hypertension, is the only vascular lesion possibly to blame which has not yet been proved to be a result of high blood pressure (Pickering 1943)

Experimental studies The classical experiments of Goldblatt (1934 *et seq*) proved that persistent hypertension could be induced in dogs by constricting both renal arteries; unilateral constriction failed unless the other kidney was removed. Hypertensive retinopathy and widespread arteriolar necrosis similar to malignant hypertension in man were reproduced by more severe constriction but the renal vessels distal to the clamp were spared. Similar results were obtained in rabbits by Wilson and Pickering (1937). In 1939 Wilson and Byrom succeeded in causing persistent hypertension, benign or malignant, in rats by constricting only one renal artery. The vessels in the other kidney then showed changes comparable in all respects to those seen in benign or malignant hypertension in man.

The conclusion that the difference between essential and malignant hypertension is merely one of degree is supported by the occasional development of malignant changes in practically all varieties of hypertension including paroxysmal transient and renal hypertension; moreover in the early malignant stage renal biopsy usually reveals no evidence of arteriolar necrosis, indicating that this is not an essential part of the picture but merely a late consequence (Castleman and Smithwick 1943).

Biochemical hypothesis concerning the cause of hypertension Experimental hypertension of the kind just described is believed to depend upon the liberation of an excess of renin by the ischaemic kidney. Renin combines with an enzyme, hypertensinogen, which is a normal constituent of the plasma globulins, to form a pressor substance, hypertensin or angiotonin (Braun Menendez *et al* 1939). Hypertensin is said to be destroyed by another enzyme, hypertensinase (Pickering 1943).

There is as yet no direct proof that essential hypertension in man is caused by this mechanism although it seems to explain renal hypertension. It may also explain rare cases of hypertension associated with atherosclerotic obstruction of one or both renal arteries (Yude 1944). It should be noted that unilateral renal disease is capable of causing hypertension in man in other words man behaves like the rat in this respect not like the dog or rabbit.

Physiology of the circulation in essential hypertension In essential hypertension vasoconstriction affects chiefly the efferent glomerular arterioles of the kidney the intraglomerular pressure being raised and the cortical blood flow diminished obviously if the latter were due to vasoconstriction proximal to the glomeruli the intraglomerular pressure would be lowered. Blood appears to be diverted from the renal cortex into other channels. The classical studies of Trueta and his colleagues (1947) make it highly probable that the juxta medullary by pass provides the principal diversion. The vessels of the skin and brain are constricted more or less sufficiently to prevent an increased blood flow through these territories, on the other hand the arterioles in skeletal muscle and probably in the heart are little if at all constricted so that they may passively yield to the raised pressure and take some of the shunt. The behaviour of the splanchnic vessels remains to be investigated. The cardiac output, blood volume, and blood viscosity are normal. Vasoconstriction appears to be humoral rather than nervous in mechanism (Pickering 1943). Hypertensin causes a similar type of vasoconstriction the chief effect is on the efferent glomerular arterioles the skin is involved only to the extent of preventing secondary increase of blood flow the skeletal muscles take some of the shunt. That hypertensin is the humoral cause of essential hypertension is therefore an attractive hypothesis. Pickering remarks that the brain and heart being two of the most important organs in the body are provided with special pressor mechanisms the carotid sinus and aortic arch which respond to falling intravascular pressure by causing vasoconstriction as the nature of these organs demands that appropriate adjustments are immediately executed it is natural that the mechanism of this vasoconstriction is nervous. But the kidneys are just as vital and it would therefore harmonise with general principles if they too were provided with a pressor mechanism to insure adequate intraglomerular pressure without which filtration would cease but there is no necessity for sudden adjustments but rather for prolonged ones. A humoral mechanism would meet the requirements nicely.

Nevertheless as previously stated proof that essential hypertension in man is due to excessive liberation of renin is lacking. Transfusion experiments have failed to demonstrate a pressor agent in the venous blood of hypertensive subjects and Light and I (1939) failed to demonstrate a pressor agent in a pint of blood taken from the renal vein of a patient with malignant hypertension and transfused into a boy of nine. Even if the humoral mechanism were proved to be the renin hypertensin system

we should still be ignorant of the cause of its hyperactivity

The most promising line of investigation seems to be that recently opened up by Trueta and his colleagues at Oxford. They have shown that blood reaching the kidney has two alternative routes (1) through the glomeruli of the cortex (2) through a juxta medullary by pass. Blood may be diverted from the cortex in varying degree as a result of emotion, shock, crushing injuries, hæmorrhages, certain drugs, certain bacterial toxins, and probably by innumerable other agents. In cases of Bright's disease they have noticed degenerative changes in the juxta medullary glomeruli consistent with constant operation of the shunt. The significance of these findings will not be overlooked, particularly their suggestion that the juxta medullary by pass may act as a functional Goldblatt clamp.

CLINICAL FEATURES

PAROXYSMAL HYPERTENSION

Paroxysmal hypertension, first described by Frankel (1886), is rare, usually occurs in youthful or early middle aged subjects of either sex, and is characterised by recurrent attacks of palpitation, headache, and vomiting, angina pectoris, or even acute pulmonary œdema. (Howard and Barker, 1937) may be associated. Abdominal compression, as occurs on stooping, may provoke an attack, but usually there is no obvious precipitating cause. During the crisis, which may last for minutes or hours, the blood pressure (systolic and diastolic) is extremely high, most of the skin is cold, pale and mottled, but the forehead, face and neck may be flushed. Sweating and trembling may follow. Between attacks the patient is usually well, but persistent hypertension, occasionally malignant, develops sooner or later in the majority (Green, 1946).

A mass about the size of an orange may be felt in the abdomen in one third of the cases, or may be demonstrated by simple skiagrams, pyelograms, or other radiological methods. The adrenal medullary tumour is commonly unilateral and benign. There is usually a considerable excess of circulating adrenaline or nor adrenaline all the time, and in attacks there may be a thousand times the normal quantity (Mackenzie, 1944). The electrocardiogram may show the usual pattern associated with persistent hypertension, or it may show evidence of acute left ventricular stress during attacks—*inversion of the T wave in leads facing the surface of the left ventricle*.

Death may result from cerebral hæmorrhage, acute pulmonary œdema, or ventricular fibrillation.

Following the demonstration by Clerc and Sterne (1957) that a synthetic benzodioxan (diethyl aminoethyl benzodioxan) in oral doses of 0.05 G, six hourly, relieved all symptoms immediately and prevented further attacks, the administration of this substance has been used as a diagnostic test for the condition (Cahill, 1948).

The intravenous injection of 0.025 mg of histamine or of 300 mg of tetraethylammonium bromide is also helpful in diagnosis for in cases of phaeochromocytoma both raise the blood pressure (La Due *et al*, 1948)

Treatment is surgical and may be entirely successful but the operative mortality is about 30 per cent (Mackenth 1944) The chief dangers are extreme hyper adrenalism during manipulation of the tumour, and a profound drop in blood pressure following its removal

TRANSIENT HYPERTENSION

The clinical features of acute nephritis and toxæmia of pregnancy are beyond the scope of this work and their effect upon the heart is discussed elsewhere (page 327)

PERSISTENT HYPERTENSION

It is doubtful whether any symptoms can be ascribed to high blood pressure itself Certainly the majority of cases are discovered accidentally or by reason of complications Headaches fatigue dizziness difficulty in concentration and palpitations are commonly due to anxiety whether the blood pressure is raised or not Redistribution of blood due to selective vasoconstriction may however determine the behaviour of two variables It was stated previously that vasoconstriction in skin and brain was more or less sufficient to prevent an increase of blood flow through these territories as a result of raised pressure the words more or less may now be amplified Thus more cutaneous vasoconstriction may be responsible for the pale hypertensive less for the red more cerebral vasoconstriction may be responsible for dizziness failing memory and for general mental deterioration less for headache The more important symptoms associated with hypertension are due to cardiac renal or cerebral complications and will be discussed later

The blood pressure is necessarily raised a diagnosis of previous persistent hypertension when the blood pressure is found to be normal is nearly always wrong unless there is severe hæmorrhage shock massive pulmonary embolism or myocardial infarction It is customary to recognise four grades of hypertension according to the level of the diastolic pressure between 90 and 110 mm Hg is considered mild 110 to 130 moderate 130 to 150 severe above 150 gross The systolic pressure may be at any level between 150 and 300 mm Hg and may modify the grade accordingly With mild hypertension it is usually between 150 and 200 with moderate hypertension between 180 and 230 with severe between 210 and 260 with gross between 240 and 300 Essential and nephritic hypertension may be of any grade malignant hypertension is always severe or gross

The pulse is firm and varies considerably in amplitude from case to case In the more severe grades it is apt to be small in those with marked atherosclerosis large Hard tortuous or calcified peripheral arteries indicate atherosclerosis or Monckeberg's sclerosis not hypertension—although they

may be associated. In the latter event one or other carotid usually the right may be knicked and then mistaken for an aneurysm or carotid pulsation may be so increased in amplitude as to suggest aortic incompetence. A diminished and delayed femoral pulse associated with absent dorsalis pedis and posterior tibial pulses indicates coarctation of the aorta. Pulsus alternans (page 166) may occur in severe cases and is usually associated with heart failure.

Retinoscopy may reveal arterial thickening, hæmorrhages, exudates or papilloedema and should never be omitted. There are five signs of arterial thickening: (1) increased tortuosity; (2) notching, pinching or S-shaped bending of veins at arterio-venous crossings; (3) uniform or irregular narrowing of the arterial blood streams owing to reduction in the diameter of the vascular lumina; (4) white arterial fringes or thin white lines bordering the red arterial streams representing the thickened white walls of the arteries themselves—they are rarely seen in more than one or two places and then only for a short distance usually on a bend; (5) the silver wire artery in which a single white streak representing a grossly thickened artery with an obliterated lumen replaces the red column of blood. This is rare and usually signifies thrombosis. Occasionally the distal part of such an artery may be patent due to the development of a collateral circulation.

By far the most important of these signs is narrowing of the arterial lumen. Normally the apparent width of a retinal artery compared with its accompanying vein is as 5 : 5 or 4 : 5. When the artery is thickened this ratio is decreased and may be about 3 : 5 or less. There is no better way of expressing the average calibre of the retinal arteries than by giving the approximate arterio-venous ratio.

In benign hypertension it is rare to find more than notching of veins and narrowing of the arterial lumina. White arterial fringes and obliteration of the lumen usually mean nephritic or malignant hypertension.

It should perhaps be added that the appearance of the fundal vessels gives little indication of the state of the cerebral vessels; the risk of stroke cannot be assessed from retinoscopy.

Retinal hæmorrhage may be superficial when it is linear or fan-shaped in appearance or deep when it resembles a rounded smudge. Both kinds may be seen in *hypertensive retinopathy* but the former is more common. Hæmorrhages are rare in essential hypertension and when present are usually minute. They are not uncommon in nephritic hypertension and almost invariable sooner or later in the malignant type.

Retinal exudates are of four distinct types: (1) large hæmorrhages sometimes reveal eccentric soft white cores which may persist after absorption of the blood; (2) soft fleecy patches scattered indiscriminately over the retina are characteristic of malignant hypertension; (3) complete or incomplete star patterns composed of hard whitish particles or dots radiating from the macula may be seen in chronic nephritic or in malignant hyper-

tension (4) in diabetes mellitus the exudate is waxy sharply cut, and scattered resembling pale yellow confetti. Small areas of retinal degeneration in old people should not be confused with exudates.

When papilloedema is added to the signs of hypertensive retinopathy already described, malignant hypertension should be diagnosed. Conversely malignant hypertension should rarely be considered in the absence of papilloedema.

Although chronic nephritis may be responsible little is lost by making the other diagnosis for if there is papilloedema the course of the disease will certainly be malignant if acute nephritis and toxæmia of pregnancy can be excluded. The appearances may be distinguished from those of cerebral tumour by the arterial changes by a macular star figure or by exudates independent of hæmorrhages.

The mechanism of hypertensive retinopathy is obscure. Hæmorrhages can hardly be due to rupture of minute vessels subjected to high pressure for the capillary pressure is normal in hypertension (Ellis and Weiss 1929-30) and in any case healthy capillaries can withstand astonishingly high pressures. Papilloedema is usually associated with a high cerebro spinal fluid pressure but not invariably; moreover higher C.S.F. pressures are found without papilloedema in cases of superior vena cava obstruction. Occasionally progressive blindness occurs.

Examination of the heart usually reveals some degree of left ventricular hypertrophy. The apex beat becomes displaced to the left and downwards the cardiac impulse becomes heaving in quality and unusually easy to feel. It is quite different from the short sharp thrust of the over acting heart for it is a quiet unhurried action giving the impression of great strength. The dynamic quality of the former may be compared with the first few strokes of a racing crew galvanised into urgent action by the sound of the starting signal the heaving impulse of left ventricular hypertrophy to the powerful steady drive maintained by the crew when it has settled down to a long hard struggle. If with due care the apex beat cannot be located left ventricular hypertrophy is unlikely even in obese subjects unless masked by emphysema.

Presystolic gallop rhythm is common with severe hypertension especially but not necessarily when complicated by left ventricular failure. The second sound at the base is accentuated and high pitched. Functional aortic incompetence is not uncommon and may be associated with diastolic pressures of 130 to 170 mm Hg. in other words it may not affect the circulatory dynamics. It is due to dilatation of the aortic ring and may be compared with functional pulmonary incompetence in mitral stenosis and atrial septal defect. Pulsus alternans may sometimes be heard especially if there is a mitral systolic murmur (Levine 1948).

Auricular fibrillation is found in about 7.5 per cent of unselected hypertensive patients (Rothstadt 1938) and may precipitate congestive heart failure. At first and particularly if untreated it may be paroxysmal but as

a rule it soon becomes persistent especially under the influence of digitalis and in elderly subjects. Permanent auricular fibrillation is less troublesome than paroxysmal and tends to protect the individual from paroxysmal cardiac dyspnoea and acute pulmonary oedema. Other rhythm changes are relatively rare but include auricular flutter, paroxysmal tachycardia and all degrees of heart block.

Limitation of cardiac reserve is indicated by undue breathlessness on exertion and by poor responses to effort tolerance tests. Left ventricular failure develops sooner or later in the majority of those who survive the other hazards of hypertension and may be recognised by a history of orthopnoea, paroxysmal cardiac dyspnoea or pulmonary oedema and by finding persistent rales at the lung bases, a diminished vital capacity and lung volume, prolongation of the crude pulmonary circulation time and exaggeration of the pulmonary vascular shadows as described on pages 158 to 163.

Congestive heart failure with elevation of the venous pressure, hepatic distension and dependent oedema follows left ventricular failure in practically all cases that survive other risks. Not infrequently patients with hypertensive heart disease develop congestive heart failure without previous orthopnoea and paroxysmal cardiac dyspnoea. No satisfactory explanation for the behaviour of these cases has yet been offered. Catheter studies have proved that the right ventricular pressure is commonly normal in hypertension and that although it rises in left ventricular failure the levels reached do not seem high enough to be responsible for right ventricular failure as much higher pressures may be found in pulmonary stenosis and patent ductus without embarrassment. The suggestion that the right ventricle is partly obstructed by displacement of the interventricular septum (Bernheim's Syndrome) lacks proof but necropsy evidence is suggestive (East and Bain 1949). Functional pulmonary stenosis was disproved in one case of the author's by means of cardiac catheterisation.

The cardiac output is low in hypertensive congestive heart failure but may be near normal at rest in left ventricular failure; moreover paroxysmal cardiac dyspnoea may occur as the output rises (page 159).

The size of the heart in hypertension bears a close relationship to the duration of heart failure: it is larger in essential hypertension when failure has been protracted, least enlarged in chronic nephritic hypertension when death is due to renal failure or in those who die from apoplexy or from other non-cardiac causes (Harrison and Wood 1949). Again serial skiagrams may show little alteration in the manifest size of the heart for long periods in essential hypertension yet gross enlargement may develop rapidly when failure occurs. This is not merely a matter of cardiac dilatation because heart weights show similar correlation. Slight to moderate left ventricular hypertrophy probably results from hypertension alone according to its degree and duration but gross enlargement which usually involves the right ventricle as well as the left is always due to protracted failure.

Moderate hypertrophy should be regarded as a compensatory change of structure which is beneficial it helps the heart to perform more work (Dieckhoff 1936)

Electrocardiography provides the most accurate means by which the degree of left ventricular enlargement and stress may be assessed. Leads facing the surface of the left ventricle such as V_5 and V_6 show high voltage and slightly widened R waves with depressed R-T segments and inverted T waves (fig 15 01). This fundamental pattern is reflected in right

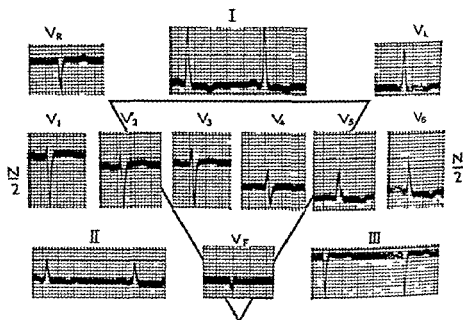


Fig 15 01—Electrocardiogram in a case of hypertensive heart disease (see text). The heart is electrically horizontal.

ventricular surface leads such as V_1 and V_2 as very small R waves and deep S waves the S-T segment being elevated and the T wave invariably upright. The heart is usually electrically horizontal, left ventricular surface potentials being transmitted to the left arm, right ventricular surface potentials to the left leg. Lead V_L then resembles V_5 and V_6 , lead V_F resembles V_1 . Standard limb leads therefore show left axis deviation, lead 1 looking like V_L and V_5 or 6, lead 3 like V_F and V_1 .

When the heart is rotated clockwise on its longitudinal axis (viewed from below) the anterior part of the inter-ventricular septum is displaced to the left and the transition zone becomes V_4 or even V_5 (fig 15 02). When the heart is rotated anti-clockwise the transition zone moves to the right and QR complexes or dominant R waves with inverted T waves may be found as far across as V_2 (fig 15 03).

When the heart is electrically vertical, left ventricular surface potentials are transmitted to the left leg, right ventricular surface potentials to the left

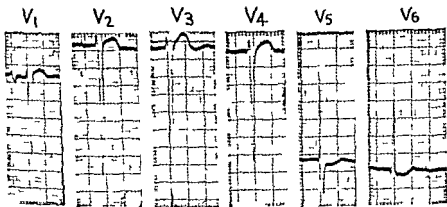


Fig 1302—Electrocardiogram in a case of hypertensive heart disease with clockwise rotation about the longitudinal axis the transition zone is shifted to the left

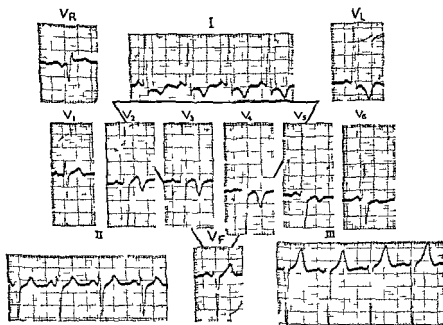


Fig 1300—Electrocardiogram in a case of hypertensive heart disease with anti-clockwise rotation about the longitudinal axis the transition zone is shifted to the right

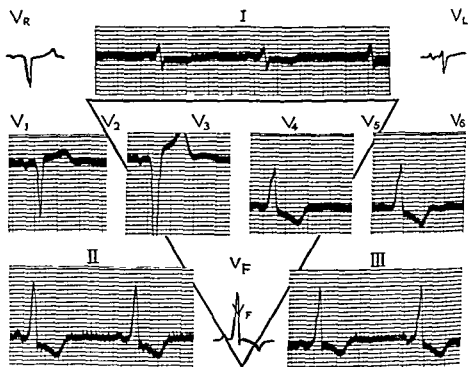


Fig 15 04—Electrocardiogram in a case of hypertensive heart disease. The heart is electrically vertical (see text)

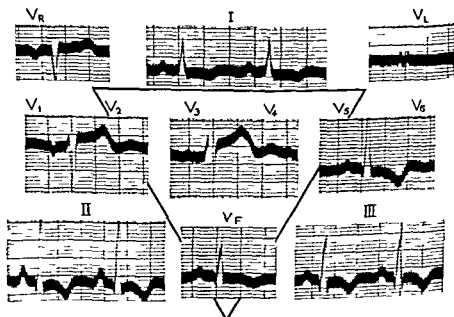


Fig 15 05—Electrocardiogram showing concordant left ventricular preponderance due to a semi vertical position of the heart

arm. Lead VF then shows the tall R wave and inverted I whilst lead VL has a prominent S wave. Standard leads may then show right axis deviation with inversion of the T wave in leads 2 and 3 (fig. 15 04).

Concordant left ventricular preponderance in standard leads (fig. 15 05) is due to a semi vertical electrical position of the heart. Left ventricular surface potentials are transmitted to the left leg and standard leads show high voltage R waves and inversion of the T wave in all leads.

The higher and wider the R wave in lead $\text{V}_5\text{-V}_6$ and the deeper the S wave in lead V_1 the bigger the left ventricle. The pattern may be distinguished from left bundle branch block by the presence of Q in lead V_6 . The cause of the R-T segment depression and the T wave inversion is less well understood; these changes may be associated with acute left ventricular stress without hypertrophy of the muscle although they usually result from both coronary disease is not responsible.

X rays reveal left ventricular enlargement fairly well but accurate measurement is difficult in obese subjects. The left border of the heart is not only displaced to the left but is denser and more rounded than usual and may sink deeply into the shadow of the diaphragm (fig. 15 06a) whilst the point of opposing movement is displaced upwards. In the second oblique position the patient often has to be turned farther to the right in order to prevent the shadow of the left ventricle overlapping the spine; the increased bulk of the left ventricle is usually obvious (fig. 15 06b).

Hypertension also leads to unfolding of the aortic arch. The ascending limb curves more forward and to the right, the descending more backward and to the left. In the antero posterior view the aorta may thus appear widened but it is only because the two limbs throw adjacent instead of superimposed shadows (fig. 15 07a). Unfolding is best seen in the left anterior oblique position especially with barium in the œsophagus which is deflected back with it (fig. 15 07b); the abrupt angulation so caused occasionally produces dysphagia. In the first oblique position the œsophageal deflection may be almost as conspicuous (fig. 15 07c). In this view backward displacement of the œsophagus at left auricular level may be due to enlargement of the base of the left ventricle (fig. 15 08).

The combination of left ventricular enlargement and unfolding of the aorta presents the characteristic appearance of two ovals set at right angles; another descriptive term is 'boot shaped' (this should not be confused with the *cœur en sabot* which compares the turned up toe of the heart in Fallot's tetralogy with that of the wooden shoe commonly worn by Dutch peasants).

Angina pectoris occurs in 5 to 10 per cent of cases and may be due to associated coronary atherosclerosis or to relative coronary insufficiency. Pain may be typical or it may tend to last longer than usual even up to an hour or so depending particularly upon transient rises of blood pressure such as occur for example in paroxysmal hypertension, strong emotion, e.g. fear or anger and exposure to cold may provoke such an attack.



(a) Antero posterior view the apex of the left ventricle is buried in the diaphragm



(b) Angiocardiogram in the second oblique position

Fig 15 06—Hypertensive heart disease showing left ventricular enlargement



(a) Anteroposterior view



(b) Left anterior oblique position

Fig 15 07—Skiagram of a case of hypertensive heart disease showing unfolding of the aorta at h



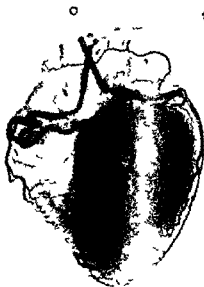
Fig 15 07 (c)—Right anterior oblique position



Fig 15 08—Right anterior oblique view of a case of hypertensive heart disease showing backward displacement of the esophagus at left vertebral level



(a)



(b)

Fig 109—Comparison of the coronary systems in a normal (a) and a hypertensive heart (b). The coronary vessels have been injected with a radio opaque gel (see text)



(a)



(b)

Fig 110—Coronary system in two cases of hypertensive heart disease with mitral regurgitation.
(a) Showing occlusion of coronary atherosclerosis (mixed type).
(b) Showing failure of the primary vessels to mark the heart.

It will be remembered that the coronary blood flow depend upon the mean blood pressure and upon the state of the coronary arteries. During systole the large extra mural coronary vessels dilate forming a tense elastic reservoir the outflow being sealed by the intramural pressure. The higher the systolic pressure the greater this elastic reservoir. As the ventricles relax blood flows through the intra mural branches influenced not only by the aortic diastolic pressure but also by the elastic recoil of the superficial coronary arteries.

Autopsy studies indicate that the coronary blood flow in essential hypertension is considerably increased. In figs 15 09a and b the coronary systems of a normal and of a hypertensive heart are compared. The vessels have been injected with a radio opaque substance at the calculated mean pressure and skiagrams have been taken at a fixed distance so that comparative measurements are valid. The large and luxuriant coronary tree of the hypertensive case is typical of the series studied (Harrison and Wood 1949). It is probable that the coronary flow behaves like the blood flow through skeletal muscle and is usually increased in all forms of hypertension. In cases of angina however skiagrams of the injected coronary vessels show either occlusive atherosclerosis (fig 15 10a) or a meagre coronary system which has failed to enlarge (fig 15 10b).

Renal behaviour varies greatly according to the type of hypertension. In the essential variety renal failure is rare and when it does occur it is late usually in patients over 70 years of age. Minor degrees of renal involvement however are common. Traces of albumin and hyaline casts are often found in the urine due to glomerular fault and diminished filtration may be revealed by inulin creatinine or urea clearance tests. Tubular re absorption may be impaired resulting in polyuria and in diminished power of urinary concentration. Nocturia may also be a feature.

In malignant hypertension there is always a fast race between renal failure cardiac failure and cerebral catastrophe. The end is sometimes a combination of all three. Nevertheless despite the early occurrence of renal failure it is rare for pronounced changes in renal function or for conspicuous urinary findings to precede the characteristic retinopathy (Wagener and Keith 1924). The converse is true of nephritic hypertension. Nephrosclerosis in malignant hypertension differs from that found in essential hypertension only in the presence of afferent glomerular arteriolar necrosis.

In chronic nephritis there is usually considerable evidence of renal damage at a time when the heart is but little enlarged and when the fundi are relatively normal. Albumin hyaline and granular casts and occasionally red cells are found in the urine inulin creatinine and urea clearance are greatly diminished the blood urea may be raised and there is commonly polyuria nocturia and failure of urinary concentration.

Cerebral manifestations occur sooner or later in about one quarter of hypertensive cases. *Cerebral hæmorrhage* is an ever present danger and

may at any time cut short the life of the patient. *Subarachnoid hæmorrhage* is by no means rare congenital deficiencies in the media or elastica of certain arteries particularly those forming the circle of Willis with or without berry aneurysm giving way to the high pressure. *Cerebral thrombosis* may also occur but depends more upon associated cerebral atherosclerosis.

Hypertensive encephalopathy is characterised by attacks of severe headache vomiting coma or convulsions lasting for hours with or without transient localising signs. Its mechanism is obscure but the customary treatment by dehydration is based on the belief that it is due to cerebral œdema and thus has limited pathological support (Scheinker, 1948). The diagnosis should never be made until cerebral or subarachnoid hæmorrhage has been excluded. Sometimes careful subsequent examination of the central nervous system reveals some abnormality indicating a local vascular lesion.

Deterioration of higher cerebral function has already been mentioned when severe it is usually due to associated atherosclerosis and ischæmia. Occasionally however multiple pin point hæmorrhages scattered widely throughout the frontal lobes are found at autopsy and provide adequate explanation for dementia.

Hæmorrhages elsewhere are not uncommon and include epistaxis hæmoptysis and hæmatemesis. Whilst some local predisposing factor would seem probable nothing significant is usually found. Clinical diagnosis in such cases may be obscure at first for hypertension may not be recognised owing to the fall of blood pressure which accompanies the hæmorrhage. Moreover when hæmodilution is slow so that the hæmoglobin or hæmatocrit level is but little reduced the apparently normal blood pressure may lead to gross error of judgment concerning the size of the hæmorrhage. Routine examination of the ocular fundi tends to prevent such mistakes.

COURSE AND PROGNOSIS

(of persistent hypertension)

Perhaps the best follow up studies in the literature are those by Janeway (1913) Blackford Bowers and Baker (1930) and Bechgaard (1946). Janeway found that one half of 458 patients were dead within five years and three quarters within ten years of the onset of symptoms. Blackford Bowers and Baker reported a 50 per cent mortality (70 per cent of the men 39 per cent of the women) amongst 222 cases within five to eleven years. Of Bechgaard's 1 000 patients 41 per cent of the men and 22·4 per cent of the women were dead within five to ten years. The better outlook in women was emphasised in all three articles. Bechgaard found the mortality rate of hypertensive men was 2·9 times and women 1·4 times, that of the general population and was similar in all age groups (excluding renal cases).

Apart from sex the chief factors affecting prognosis include the type of hypertension the degree of retinopathy the height of the blood pressure

and the state of the heart. The natural outlook in malignant hypertension is uniformly bad: few cases surviving more than one or two years. Chronic nephritic hypertension also has a grave prognosis: the mortality rate being about nine times that of essential hypertension. This is partly because renal hypertension is often a late manifestation of chronic kidney disease—hence the frequency of a normal sized heart in this group.

Wagener and Keith (1939) correlated life expectancy with changes in the ocular fundi: they followed the course of 209 patients for five to nine years. The survival rate according to whether retinal changes were mild, moderate, severe or gross was 80 per cent, 35 per cent, 9 per cent and nil respectively. When retinopathy was gross and included papilloedema, 80 per cent died within one year.

The height of the blood pressure matters little so long as it remains within the slight or moderate grade, i.e. under 130 mm Hg diastolic, but above this level it is always serious.

Cardiac behaviour in hypertension is determined by the amount of extra work involved, and by the ability of the heart to cope with it: it is chiefly influenced by the rapidity of hypertensive development, by the size and strength of the left ventricle, and by the efficiency of the coronary blood flow. The best defence is put up by a placid patient of voluntary or enforced sedentary habits and occupation, who has a naturally strong left ventricle with a good coronary blood flow: when hypertension is neither too severe nor too sudden, under such circumstances the heart enlarges but little over the years, failure is indefinitely deferred, and the patient remains free from cardiac symptoms. The worst defence, leading to rapid failure, and perhaps to early death, occurs in an excitable individual of active physical habits and strenuous occupation, who tries to cope with a rapidly developing and extreme hypertension with an unprepared left ventricle indifferently nourished by a mean coronary system.

Evidence of any cardiac abnormality, e.g. diminished cardiac reserve, angina pectoris, enlargement or electrocardiographic changes, at once doubles or trebles the mortality rate (Bechgaard, 1946). Inversion of the T wave in left ventricular surface leads or their equivalent is particularly grave, at least 60 per cent of such cases being dead in an average of eight months from the time of its discovery (Rykert and Hepburn, 1955). Atrial fibrillation means death within two years in 80 per cent of cases (Rothstadt, 1938).

Hypertensive heart failure is characteristically left ventricular at first and limits life expectancy to about eighteen months. Systemic congestion follows sooner or later. Several congestive attacks usually occur, each responding less satisfactorily to treatment than its predecessor. The patient finally sinks into a stuporose condition with chronic venous congestion, hepatic engorgement and dependent dropsy: the blood pressure falls, Cheyne Stokes breathing develops, and death comes slowly. Heart disease is responsible for death in 33 per cent (Janeway, 1913) to 55 per cent (Bell

and Clawson 1928) of hypertensive cases stroke in 7.2 per cent (Paullin *et al* 1927) to 16 per cent (Bechgaard 1946) uræmia in 10 per cent (Bechgaard 1946)

The average life expectancy in uncomplicated benign hypertension of slight or moderate grade is about fifteen years (Fahr, 1928). Obese subjects do as well or better than those with normal weight probably because their blood pressures are not as high as they seem possibly because of benefits derived from weight reduction. Spontaneous recovery occurred in 5.4 per cent of Bechgaard's series (2 per cent of the women 13 per cent of the men) but in none of those seen by Blackford Bowers and Baker. After five to ten years 58 per cent of Bechgaard's cases were free from symptoms or only slightly inconvenienced.

Only about 0.2 per cent of cases of simple high blood pressure develop malignant hypertension but 8 per cent of cases of chronic pyelonephritis do so.

TREATMENT

It must be said at once that as yet there is no satisfactory treatment for essential or for malignant hypertension when nephritic hypertension is due to a unilateral lesion such as chronic pyelonephritis nephrectomy may be curative but otherwise it can be little influenced. The hypertension of Cushing's syndrome responds to removal of the offending tumour and that due to coarctation of the aorta to surgical repair. When hypertension is associated with unilateral renal disease a causal relationship cannot always be assumed before advising nephrectomy it is well to make sure that neither parent was hypertensive (Platt 1947). Normal renal function is also a necessary condition for successful nephrectomy for severe hypertension may have so damaged the vessels of the originally healthy kidney as to have made it ischæmic and so to have established a vicious circle (Wilson and Byron 1941).

For essential hypertension there are four main lines of treatment (1) conservative (2) the low sodium or rice diet (3) thiocyanates (4) lumbo-dorsal sympathectomy.

Conservative. When the grade of hypertension is mild or moderate and when the prognosis is judged to be good on criteria previously outlined radical medical or surgical treatment is hardly justified but this does not mean that nothing else need be done. Conservative treatment seeks to correct adverse factors and to prevent complications or deterioration.

If circumstances permit it is a good plan to begin treatment by putting the patient to bed and to keep him there until the blood pressure has reached a static level. Symptoms usually disappear quickly and the patient gains confidence. During this time renal function including pyelography may be fully and conveniently investigated also the reaction of the blood pressure to bed rest gives useful diagnostic and prognostic information.

innocent labile types falling quickly to normal nephritic and malignant hypertension responding least

Patients should then be advised to live at a lower tempo they should learn to refuse extra commitments and gradually to relinquish the least important or most irksome of those they already have they should keep all Saturday and Sunday free for relaxation should have at least nine hours rest in bed every night and should insist on proper holidays each year preferably six weeks Long working hours heavy mental or physical stress and the general rush hurry and struggle of modern life must be avoided or reduced Occupation may require modification but it is rarely practicable to change it radically for the patients are generally too old Sudden effort especially in the cold or after a heavy meal should be avoided straining at stool should be prevented by regular habits and if necessary by the use of liquid paraffin

Mental relaxation may be impossible without sedatives or psychiatric help Phenobarbitone $\frac{1}{2}$ to 1 grain (32 to 64 mg) t d s may be prescribed at times of unavoidable anxiety alternated with potassium bromide 5 to 10 grains (0.32 to 0.65 G) t d s Psychiatric help is invaluable not necessarily from a psychiatrist but by any experienced physician with the requisite knowledge Many of the symptoms ascribed to hypertension are more often due to anxiety moreover hypertensives usually have hyper reactions to anxiety in the sense that their blood pressures rise unduly (Hines 1940) Details of psychiatric treatment are given on page 543

Symptoms attributed to hypertension at the menopause may respond to stilboestrol 1 to 3 mg daily although the blood pressure does not fall an associated anxiety state is also common at this time

Obese patients tend to do well on a weight reducing diet This is not merely because they lose weight but because small meals are beneficial to hypertensives One day's bed rest with semi starvation per week diet then being limited to fresh fruit fruit juice and water only may be most helpful or such a regime may be instituted at less frequent intervals when the patient feels the need of it

The use of harmless blood pressure reducing drugs other than sedatives is no part of conservative treatment for none substantiates the claim made for it (Evans and Loughnan 1939) Thiocyanate will be considered later

Venesection has been advocated in the past and is still practised from time to time It is only justified in phlethoric cases associated with polycythaemia In essential hypertension its effect is fleeting the blood pressure often regaining its previous level within twenty four hours In malignant hypertension and in chronic nephritis venesection is contra indicated for some degree of anaemia is usually present in both conditions

Low sodium diet A low sodium diet consisting of soya beans peanut flour cooked potatoes rice starch and gelatine lowered the blood pressure in experimental hypertension in rats (Grollman and Harrison 1945) Subsequent clinical studies have shown that a similar diet may reduce the

blood pressure substantially in essential human hypertension (Grollman 1945). A low sodium diet is described in detail on page 185. The rice diet (Kempner 1946) consists of rice cooked in unsalted water fruit in any form and sugar fruit juice and fluids are given freely. This contains less than 0.5 G of sodium per day.

The blood pressure is said to fall significantly (average 45/25) in about 60 per cent of cases and there is usually a loss in weight of about 7 lb (3 kg). As soon as such benefit is demonstrated the diet may be modified by adding meat vegetables and other foods listed on page 186. It is not practicable to maintain the initial rigid diet for long nor to maintain the daily sodium intake below 1 G for more than a few weeks.

Treatment of this kind or by semi starvation is useful to check hypertensive crises or to lower the blood pressure quickly when it is found to be dangerously high. The value of the modified diet in maintaining pressures at lower levels is less well established.

Thiocyanates Thiocyanate was originally introduced as a hypotensive agent by Treupel and Edinger (1900) but gained no immediate favour in view of the difficulty experienced in avoiding serious toxic symptoms. Considerable interest has been taken in the drug however since Barker (1936) showed that the dose could be properly controlled if the thiocyanate blood level was estimated weekly. The normal serum thiocyanate ranges between 0 and 2.77 mg per cent and is not altered in hypertension (Connell Wharton and Robinson 1946). Levels above 15 mg per cent are dangerous and those between 12 and 15 mg per cent are risky. Toxic symptoms include weakness anorexia indigestion nausea vomiting limb pains impotence purpura dermatitis goitre thrombophlebitis mental lethargy and confusion. In fatal cases dysarthria verbal aphasia convulsions hallucinations delirium and mania have usually preceded death by three to nineteen days (Del Solar *et al.* 1945). Progressive anaemia and emaciation have been attributed to chronic poisoning after five to ten years continuous therapy (Wald Lindberg and Barker 1939).

The potassium salt is given by mouth in initial doses of 2 to 3 grains (0.13 to 0.2 G) three times daily after meals. The serum thiocyanate is measured on the seventh day and then at weekly intervals subsequent dosage being regulated as follows.

<i>Thiocyanate level</i>	<i>Dosage recommended</i>
Under 5 mg per cent	2 to 3 grains (0.13 to 0.2 G) t d s
5 to 7	1.5 grains (0.1 G) t d s
7 to 10	1 grain (0.064 G) t d s
Over 10	Stop drug for one week

The lowest blood level compatible with a satisfactory hypotensive effect should be maintained for three to six months. Further courses may be given as desired.

The drug is said to be unsafe in patients who are over 60 years old who have had cerebral or other thrombosis or who have poor renal function but Watkinson and Evans (1947) observed no ill effect in fifteen patients over 60 nor in sixteen cases of malignant or chronic nephritic hypertension.

Thiocyanates have been particularly recommended for labile hypertensives who complain of headache and giddiness (Hines 1946) but they have also been used for severe or gross cases unsuitable for lumbo dorsal sympathectomy and as an adjunct to surgical treatment.

Clinical benefit associated with a significant fall of blood pressure has been claimed in about 60 per cent of cases (Watkinson and Evans 1947). This figure is not impressive when it is recollected that Bechgaard found that 58 per cent of 1 000 persistent hypertensives did well without treatment. Carefully controlled observations such as those by Rusken and McKinley (1947) are more convincing and throw considerable doubt on the efficacy of thiocyanates. It is well to remember that Pauli (1903) who is usually credited with introducing thiocyanate for the treatment of hypertension actually used the drug in the hope that it would prove superior to bromide in allaying anxiety symptoms and reported singular success in this respect. The effect of sedation on hypertension is well known and if thiocyanate merely acts in this way it should be abandoned in favour of less toxic substances.

Lumbo dorsal sympathectomy. In recent years numerous attempts have been made to lower the blood pressure by surgical means. The only operation which has proved eminently successful is nephrectomy in those relatively rare cases in which hypertension is due to unilateral renal disease such as chronic pyelonephritis. Of other surgical measures the best known is lumbo dorsal sympathectomy as elaborated by Smithwick (1940). This consists of bilateral resection of the whole sympathetic chain from D8 to L2 including preganglionic fibres, ganglia and splanchnic nerves. It has proved superior to Adson's subdiaphragmatic splanchnicectomy with resection of the first and second lumbar ganglia (Allen and Adson 1940) and to Peet's supradiaphragmatic splanchnicectomy with lower dorsal ganglionectomy (Peet, Woods and Braden 1940). The object is to release as much vasoconstrictor tone as possible to prevent renal cortical vasoconstriction to produce postural hypotension and of course to lower the basal blood pressure if possible. With these aims there has been an increasing tendency to extend Smithwick's operation and a number of surgeons e.g. Grimson (1947) and Boyd (1948) now favour either total or subtotal paravertebral sympathectomy, splanchnicectomy and coeliac ganglionectomy.

The results of these various procedures have been fair. The operative mortality has averaged 3.9 per cent but about 25 per cent have died during the period of post operative observation. There is no doubt that headache, dizziness and other symptoms may be alleviated, that the blood pressure may be lowered, the electrocardiogram improved, the heart size reduced.

and that retinopathy may be diminished by such means objective improvement of one kind or another has been demonstrable in about 60 per cent of cases (Smithwick 1944) The results published by different workers are usually difficult to compare because cases are apt to be selected on different criteria

Early persistent hypertension of moderate grade before degenerative vascular changes have set in should respond best but the outlook is then sufficiently good to restrain most physicians from advising surgical treatment yet if left until serious complications have arisen it may be too late This is the crux of the problem The best compromise may be to advise lumbo dorsal or more extensive sympathectomy in relatively young hypertensive subjects (under 50) whose pressures are known to be rising or whose casual diastolic pressures tend to exceed 130 mm Hg after proper medical measures have been instituted

It is not easy to predict which cases will do well The sedation test is perhaps as good as any it consists of giving sodium amytal 3 grains (0.2 G) hourly for three doses while the blood pressure is recorded every hour The greater the drop the more likely is the operation to be successful

Surgical treatment should not be undertaken if the patient is over 50 years old if renal function is seriously impaired if there is congestive heart failure or if there has been myocardial infarction Angina pectoris is no deterrent nor is a previous stroke early left ventricular failure that responds well to medical treatment is likewise no contraindication Obese subjects are technically more difficult and are probably better avoided not only on that account but also because they may do fairly well if treated medically Women are often preferred to men because they usually do better but it should be remembered that the prognosis is twice as good in women if untreated

REFERENCES

- Allen F V and Adson A W (1940) Treatment of hypertension medical versus surgical *Ann intern Med* 14 288
 Ayman A and Goldshine A D (1939) The breath holding test a simple standard stimulus of blood pressure *Arch intern Med* 63 899
 Ayman D (1934) Heredity in arteriolar (essential) hypertension a clinical study of the blood pressure of 1 524 members of 277 families *Ibid* 53 792
 Barker M H (1936) Blood cyanates in treatment of hypertension *J Amer med Ass* 106 762
 Bechgaard P (1946) Arterial hypertension A follow up study of one thousand hypertonics *Acta med Scand Supp* 172
 Beer E King F H and Prinzmetal M (1937) Pheochromocytoma with demonstration of pressor (adrenaline) substance in the blood pre operative during hypertensive crises *Ann Surg* 106 85
 Bell E T and Clawson B J (1928) Primary (essential) hypertension *Arch Path* 5 939
 Blackford J M Dowers J M and Baker J W (1930) Follow up study of hypertension *J Amer med Ass* 94 328

- Boyd A M (1948) Discussion of the surgical treatment of hypertension *Proc Roy Soc Med* 41 370
- Braasch W F Waters W and Hammer H J (1940) Hypertension and the surgical kidney *J Amer med Ass* 115 1837
- Braun Menendez E (1939) The blood pressure raising substance in the blood of ischaemic kidneys *Rev Soc Argent de Biol* 15 420
- Cahill G F (1948) Pheochromocytomas *J Amer med Ass* 138 180
- Castleman B and Smithwick R H (1943) The relation of vascular disease to the hypertensive state Based on a study of renal biopsies from one hundred hypertensive patients *Ibid* 121 1256
- Cavelti P A and Cavelti E S (1945) Studies on the pathogenesis of glomerulonephritis I Production of auto antibodies to kidney in experimental animals *Arch Path* 39 148
- Clerc A and Sterne J (1937) A case of repeated anginal crises with paroxysmal hypertension and vasomotor disturbances a record of medical treatment two surgical interventions and of the efficiency of a synthetic sympathicolytic drug *Bull et Mem Soc med d hop d Paris* 53 562
- Connell W F Wharton G K and Robinson C E (1946) The relationship of blood pressure and serum thiocyanate *Amer J med Sc* 211 74
- Dieckhoff J (1936) Leistungsfähigkeit aortenklappeninsuffizienter Herzen ohne und mit Hypertrophie im Herz Lungen Präparat (Nebst Digitalisierungseffekten) *Arch f Exper Path u Pharmacol* 182 268
- Del Solar A V Dussailant G G Brodsky M B and Rodriguez G C (1945) Fatal poisoning from potassium thiocyanate used in treatment of hypertension Report of a case and review of the literature *Arch intern Med* 75 241
- Donnison C P (1929) Blood pressure in African natives its bearing upon aetiology of hyperpiesia and arteriosclerosis *Lancet* 1 6
- East T and Bain C (1949) Right ventricular stenosis (Bernheim's syndrome) *Brit Heart J* 11 145
- Ehrstrom R (1918) Nefrosklerosen *Finska Lak sällsk handl Helsingfors* 60 365
- Ellis L B and Weiss S (1929 30) Measurement of capillary pressure under natural conditions and after arteriolar dilatation in normal subjects and in patients with arterial hypertension and with arteriosclerosis *J clin Invest* 8 47
- Evans W and Loughnan O (1939) The drug treatment of hyperpiesia *Brit Heart J* 1 199
- Fahr G (1938) Hypertension heart *Amer J med Sc* 175 453
- Fishberg A M (1939) Hypertension and nephritis London 4th ed
- Frankel F (1886) Ein Fall von doppelertigem völlig latent verlaufenen Nebennierentumor und gleichzeitiger Nephritis mit Veränderungen am circulatorischen apparat und Retinitis *Nurichous Arch* 103 244
- Goldblatt H (1937) Studies in experimental hypertension V The pathogenesis of experimental hypertension due to renal ischaemia *Ann intern Med* 11 69 — (1938) Studies on experimental hypertension VII The production of the malignant phase of hypertension *J exper Med* 67 809 — (1948) The renal origin of hypertension Springfield Illinois — Lynch J R F
- Hazal F and Summerville W W (1934) Studies on experimental hypertension I The production of persistent elevation of systolic blood pressure by means of renal ischaemia *Ibid* 59 347
- Golden A Dexter L and Weiss S (1943) Vascular disease following toxæmia of pregnancy *Arch intern Med* 72 301
- Grant R T (1935) Observations on the after histories of men suffering from the effort syndrome *Heart* 12 121
- Green D M (1946) Pheochromocytoma and chronic hypertension *J Amer med Ass* 131 1260
- Grimson K S (1947) The surgical treatment of hypertension *Advances intern Med* 2 173

Grollman A (1945) Sodium restriction in diet for hypertension *J Amer med Ass* 129 533 — and Harrison T R (1945) Effect of rigid sodium restriction on blood pressure and survival of hypertensive rats *Proc Soc exper Biol and Med* 60 52

Harrison C V and Wood P H (1949) Hypertensive and ischaemic heart disease *Brit Heart J* 11 205

Harris H A (1927) Vascular diseases and sympathetic system *Brit med J* 1 789

Hines E A (1940) The significance of vascular hyper reaction as measured by the cold pressor test *Amer Heart J* 19 408 — (1940) The hereditary factor and subsequent development of hypertension *Proc Mayo Clin* 15 145

Hines E A Jr (1946) Thiocyanates in treatment of hypertensive disease *M Clin N Amer* 30 869

Howard J E and Barker W H (1937) Paroxysmal hypertension and other clinical manifestations associated with benign chromaffin cell tumors (pheochromocytomata) *Bull Johns Hopk Hosp* 61 371

Janeway T C (1913) A clinical study of hypertensive vascular disease *Arch intern Med* 12 755

Kempner W (1946) Some effects of rice diet treatment of kidney disease and hypertension *Bull N Y Acad Med* 22 358 — (1948) Treatment of hypertensive vascular disease with rice diet *Amer J Med* 4 545

Kylin E (1926) Die Hypertoniekrankheiten Berlin pp 32 62

La Due J S, Murison P J and Pack G T (1948) The use of tetraethylammonium bromide as a diagnostic test for pheochromocytoma *Ann intern Med* 29 914

Levine S A (1948) Auscultation of the heart *Brit Heart J* 10 213

Light A I and Wood P H (1939) Protocol at Post graduate Medical School of London

Longcope W T and Winkenwerder W L (1933) Clinical features of the contracted kidney due to pyelonephritis *Johns Hopk Hosp Bull* 53 255

Mackenth R (1944) Adrenal sympathetic syndrome Chromaffin tissue tumour with paroxysmal hypertension *Brit Heart J* 6 1

Master A M, Marks H H and Dack S (1943) Hypertension in people over 40 *J Amer med Ass* 121 1251

Pauli W (1903) Ueber Ionenwirkungen und ihre therapeutische Verwendung *Munch Med Wchnschr* 50 153

Paulin J E, Bowcock H M and Wood R H (1927) Complications of hypertension *Amer Heart J* 2 613

Peet M M, Woods W W and Braden S (1940) The surgical treatment of hypertension *J Amer med Ass* 115 1875

Pickering G W (1939) The problem of high blood pressure in man *Brit med J* 1 1 — (1943) Circulation in arterial hypertension *Ibid* 11 31

Platt R (1947) Hypertension and unilateral kidney disease *Quart J Med* 16 143 — (1947) Heredity in hypertension *Ibid* 16 111 — (1948)

Severe hypertension in young person A study of 50 cases *Ibid* 17 83

Rothstadt L F (1938) The effect of auricular fibrillation on the course of hypertension *Med J Australia* 1 813

Rusken A and McKinley W F (1947) Comparative study of potassium thiocyanate and other drugs in the treatment of essential hypertension *Amer Heart J* 34 691

Rykert H E and Hepburn J (1935) Electrocardiographic abnormalities characteristic of certain cases of arterial hypertension *Ibid* 10 94

Rytand D A (1938) The renal factor in arterial hypertension with coarctation of the aorta *J clin Invest* 17 391

- Scheinker I M (1948) Hypertensive cerebral swelling a characteristic clinicopathologic syndrome *Ann intern Med* 28 630
- Schoen R (1930) Über die Doppelseitige chronische pyelogene Nephritis *Deutsches Arch f klin Med* 169 337
- Smithwick R H (1940) Technique for splanchnic resection for hypertension Preliminary report *Surgery* 7 1 — (1944) Surgical treatment of hypertension the effect of radical (lumbodorsal) splanchnicectomy on the hypertensive state of one hundred and fifty six patients followed one to five years *Arch Surgery* 49 180
- Treupel G and Edinger A (1900) Untersuchungen über Rhodan Verbindungen *Munch Med Wschr* 47 717
- Trueta J Barclay A E Daniel P M Franklin K J and Prichard M M L (1947) Studies of the renal circulation Oxford
- Wagener H P and Keith N M (1939) Diffuse arteriolar disease with hypertension and the associated retinal lesions *Medicine* 18 317 — — (1924) Cases of marked hypertension adequate renal function and neuroretinitis *Arch intern Med* 34 374
- Wald M H Lindberg H A and Barker M H (1939) Toxic manifestations of thiocyanates *J Amer med Ass* 112 1120
- Watkinson C and Evans G (1947) Potassium thiocyanate in the treatment of hypertension *Brit med J* 1 595
- Wilson C and Byrom F B (1939) Renal changes in malignant hypertension Experimental evidence *Lancet* 1 136 — — (1941) The vicious circle in chronic Bright's disease Experimental evidence from the hypertensive rat *Quart J Med* 10 65 — Pickering G W (1937-8) Acute arterial lesions in rabbits with experimental renal hypertension *Clin Sc* 3 343
- Wood P H (1941) Da Costa's syndrome *Brit med J* 1 767 805 845
- Yule C L (1944) Obstructive lesions of the main renal artery in relation to hypertension *Amer J med Sc* 207 394

Grollman A (1945) Sodium restriction in diet for hypertension *J Amer med Ass* 129 533 — and Harrison T R (1945) Effect of rigid sodium restriction on blood pressure and survival of hypertensive rats *Proc Soc exper Biol and Med* 60 52

Harrison C V and Wood P H (1949) Hypertensive and ischaemic heart disease *Brit Heart J* 11 205

Harris H A (1927) Vascular diseases and sympathetic system *Brit med J* 1 789

Hines E A (1940) The significance of vascular hyper reaction as measured by the cold pressor test *Amer Heart J* 19 408 — (1949) The hereditary factor and subsequent development of hypertension *Proc Mayo Clin* 15 145

Hines E A Jr (1946) Thiocyanates in treatment of hypertensive disease *W Clin N Amer* 30 869

Howard J E and Barker W H (1937) Paroxysmal hypertension and other clinical manifestations associated with benign chromaffin cell tumors (pheochromocytomata) *Bull Johns Hopk Hosp* 61 371

Janeway T C (1913) A clinical study of hypertensive vascular disease *Arch intern Med* 12 755

Jempner W (1946) Some effects of rice diet treatment of kidney disease and hypertension *Bull N Y Acad Med* 22 358 — (1948) Treatment of hypertensive vascular disease with rice diet *Amer J Med* 4 545

Kylin E (19 6) Die Hypertoniekrankheiten Berlin pp 32 62

La Due J S, Murison P J and Pack G T (1948) The use of tetra ethylammonium bromide as a diagnostic test for pheochromocytoma *Ann intern Med* 29 914

Levine S A (1948) Auscultation of the heart *Brit Heart J* 10 213

Light A L and Wood P H (1939) Protocol at Post graduate Medical School of London

Longcope W T and Winkenwerder W L (1933) Clinical features of the contracted kidney due to pyelonephritis *Johns Hopk Hosp Bull* 53 255

Mackenzie R (1944) Adrenal sympathetic syndrome Chromaffin tissue tumour with paroxysmal hypertension *Brit Heart J* 6 1

Master A M, Marks H H and Dack S (1943) Hypertension in people over 40 *J Amer med Ass* 121 1251

Pauli W (1903) Ueber Ionenwirkungen und ihre therapeutische Verwendung *Munch Med Wchschr* 50 153

Paullin J E, Bowcock H M and Wood R H (1927) Complications of hypertension *Amer Heart J* 2 613

Peet M M, Woods W W and Braden S (1940) The surgical treatment of hypertension *J Amer med Ass* 115 1875

Pickering G W (1939) The problem of high blood pressure in man *Brit med J* 1 1 — (1943) Circulation in arterial hypertension *Ibid* 11 1 31

Platt R (1947) Hypertension and unilateral kidney disease *Quart J Med* 16 143 — (1947) Heredity in hypertension *Ibid* 16 111 — (1948) Severe hypertension in young persons. A study of 50 cases *Ibid* 17 83

Rothstadt I E (1938) The effect of auricular fibrillation on the course of hypertension *Med J Australia* 1 813

Rusken A and McKinley W F (1947) Comparative study of potassium thiocyanate and other drugs in the treatment of essential hypertension *Amer Heart J* 34 691

Rykert H E and Hepburn J (1935) Electrocardiographic abnormalities characteristic of certain cases of arterial hypertension *Ibid* 10 942

Ryland D A (1938) The renal factor in arterial hypertension with coarctation of the aorta *J clin Invest* 17 391

- Scheinker I M (1948) Hypertensive cerebral swelling a characteristic clinicopathologic syndrome *Ann intern Med* 28 630
- Schoen R (1930) Über die Doppelseitige chronische pyelogene Nephritis *Deutsches Arch f klin Med* 169 337
- Smithwick R H (1940) Technique for splanchnic resection for hypertension Preliminary report *Surgery* 7 1 — (1944) Surgical treatment of hypertension the effect of radical (lumbodorsal) splanchnicectomy on the hypertensive state of one hundred and fifty six patients followed one to five years *Arch Surgery* 49 180
- Treupel G and Edinger A (1900) Untersuchungen ube Rhodan Verbindungen *Munch Med Wschr* 47 717
- Trueta J Barclay A E Daniel P M Franklin K J and Prichard M M L (1947) Studies of the renal circulation Oxford
- Wagener H P and Keith N M (1939) Diffuse arteriolar disease with hypertension and the associated retinal lesions *Medicine* 18 317 — — (1924) Cases of marked hypertension adequate renal function and neuroretinitis *Arch intern Med* 34 374
- Wald M H Lindberg H A and Barker M H (1939) Toxic manifestations of thiocyanates *J Amer med Ass* 112 1120
- Watkinson G and Evans G (1947) Potassium thiocyanate in the treatment of hypertension *Brit med J* 1 595
- Wilson C and Byrom F B (1939) Renal changes in malignant hypertension Experimental evidence *Lancet* 1 136 — — (1941) The vicious circle in chronic Bright's disease Experimental evidence from the hypertensive rat *Quart J Med* 10 65 — Pickering G W (1937-8) Acute arterial lesions in rabbits with experimental renal hypertension *Clin Sc* 3 343
- Wood P H (1941) Da Costa's syndrome *Brit med J* 1 767 805 845
- Yuile C L (1944) Obstructive lesions of the main renal artery in relation to hypertension *Amer J med Sc* 207 394

PULMONARY EMBOLISM

PULMONARY embolism may cause acute or subacute pulmonary heart disease sudden death from reflex ventricular fibrillation or cardiac standstill pulmonary infarction or no ill effects Emboli may be single or multiple infected or sterile They are usually attributable to mobile venous thrombi originating in the legs but are occasionally due to fat air foreign body or to fragments of some remote malignant tumour

THROMBO EMBOLISM

General incidence Massive pulmonary embolism is directly or chiefly responsible for about 3 per cent of all hospital deaths published figures ranging from 2 to 6.5 per cent as shown below

<i>Author</i>	<i>Number of necropsies</i>	<i>Incidence of fatal pulmonary embolism per cent</i>
Belt (1934)	567	6.5
Collins (1936)	10,940	2.07
Pilcher (1939)	2,861	4.5
Hampton and Castleman (1940)	3,500	3.5
McCartney (1945)	25,771	2.62
Crutcher (1948)	2,580	2.14

The actual incidence of clinical thrombo embolism among all hospital patients is difficult to assess but appears to be about 1 per cent in post operative cases (Nygaard *et al.* 1940-41) and is believed to be equally frequent in medical obstetrical and gynaecological wards (Belt 1939)

ETIOLOGY

Thrombo embolism results from the breaking away of a blood clot formed either in the right side of the heart or in the systemic venous system

Intracardiac thrombosis Clots may form in the heart in cases of mitral stenosis auricular fibrillation congestive failure myocardial infarction bacterial endocarditis and Fiedler's carditis Damaged endocardium slowing of the blood flow and backwater eddies in dilated chambers are important contributory factors In mitral stenosis clots are more likely to form

in the left than in the right auricle unless there is congestive heart failure. Myocardial infarction practically always affects the left ventricle and as mural thrombi are limited to the area of devitalised tissue emboli from the heart are usually systemic exceptions are associated with septal infarction. Bacterial endocarditis is only a source of pulmonary emboli when it affects the pulmonary or tricuspid valve a ventricular septal defect or a patent ductus arteriosus infarcts so produced are apt to be small and frequent the clinical course resembling subacute or chronic hæmorrhagic broncho pneumonia Fiedler's carditis is mentioned because it is sometimes complicated by mural thrombi which may break away and cause repeated hæmoptysis.

Venous thrombosis Intracardiac thrombosis however accounts for only about 10 per cent of pulmonary emboli (Belt 1939) the remainder including the majority of those associated with mitral stenosis auricular fibrillation myocardial infarction and congestive heart failure being due to venous thrombosis particularly in the legs. Such emboli are common Belt (1939) found them in 6 per cent of 1 990 consecutive necropsies and as often in medical as in surgical cases. They were directly responsible for death in 22 instances (1.1 per cent) and a contributory factor in approximately 70 (3.5 per cent).

The chief causes of venous thrombosis may be listed under three main headings

1 *Local venous injury*

- (a) Inflammatory—as in thrombophlebitis
- (b) Chemical—from the injection of irritant solutions
- (c) Traumatic—as in fractures
- (d) Infiltrative—as in cancer

2 *Slowing of the venous blood flow*

- (a) By local obstruction—as by tight bandaging immobilisation in an unfavourable posture or space filling lesions (including obesity)
- (b) As a whole—as in heart failure

3 *Increased clotting tendency of the blood*

- (a) Post operative post traumatic and puerperal states
- (b) Polycythæmia and hæmoconcentration
- (c) Associated with tissue breakdown e.g. carcinoma of the stomach myocardial infarction
- (d) Certain fevers e.g. typhoid

The mechanism of intravascular clotting is a complicated process and is not yet fully understood. Its discussion is beyond the scope of this work and the reader is referred to the excellent monograph by Nygaard (1941).

Thrombophlebitis is often said to be less dangerous than simple phlebotrombosis but this is doubtful for the former is less frequent but more

easily diagnosed whilst the latter is common but apt to be overlooked so that the percentage incidence of embolism is likely to appear higher in phlebothrombosis. Fatal pulmonary embolism follows the injection treatment of varicose veins in 0.05 per cent of cases (Westerborn 1937) and the operative treatment in 0.4 per cent (Westerborn 1937, McPheeters and Rice 1928).

Fractures particularly of the legs or pelvis may cause thrombo embolism on account of injury to veins, immobilisation and post traumatic acceleration of the clotting time. They may also give rise to fat embolism. Malignant neoplasms, especially carcinoma of the stomach, may be responsible for thrombo embolism as a result of venous infiltration, mechanical venous obstruction and shortening of the clotting time (owing to tissue necrosis). They may also give rise to malignant cellular emboli.

The most common cause of phlebothrombosis is immobilisation in bed especially in obese subjects over 40 years of age. Of 229 cases of fatal post operative pulmonary embolism, Prettin (1936) found the average weight in women was 12 kg. above normal and in men 4.2 kg. In a series at the Mayo clinic 93 per cent of fatal post operative pulmonary emboli occurred in patients over 40 years of age (Barnes 1937).

Congestive heart failure encourages phlebothrombosis because the circulation is slowed. When the cardiac output remains elevated or is less reduced than usual, as in failure from the hyperkinetic circulatory states, thrombosis is rare. Congestive failure due to mitral stenosis or to myocardial infarction is particularly dangerous. Eppinger and Kennedy (1938) found that pulmonary embolism was the direct cause of death in 6.5 per cent of 200 fatal cases of coronary thrombosis and a contributory cause in 32.7 per cent of those with congestive failure. The clotting time appears to shorten after myocardial infarction, perhaps owing to the products of tissue necrosis. The clotting time also appears to be shortened by digitalis (Massie *et al.* 1944) and by the organic mercurial diuretics (Macht, 1946).

Almost any major surgical procedure may result in thrombo embolism but abdominal and pelvic operations carry the highest embolic risk. Responsible factors include post operative reduction of the clotting time (maximum at the tenth day) and immobilisation. Child birth incurs a similar risk for similar reasons. McCartnev (1945) found that pulmonary embolism was directly responsible for 5.28 per cent of obstetrical fatalities and for 5.1 per cent of post operative deaths.

HAEMODYNAMICS AND PATHOLOGY

Acute pulmonary heart disease. Experiments in which the pulmonary arteries have been occluded in varying degree by ligature or by artificial emboli have shown that it is necessary to obstruct about 60 to 85 per cent of their total cross section before the systemic blood pressure falls or before signs of right ventricular failure can be detected and between 85 and 100 per cent before death ensues (Gibbon, Hopkinson and Churchill 1932). It

is thus possible to undertake unilateral pneumonectomy without embarrassing the circulation (Barnes 1941). In accord with these facts the majority of pulmonary emboli cause no cardiac disturbance but when a large embolus lodges at the bifurcation of the main pulmonary artery or when multiple emboli block more than two thirds of the more distal trunks the circulation is impeded and the left ventricular output falls. This is the condition known as massive pulmonary embolism its effect upon the heart is acute pulmonary heart disease. Compensatory adjustments include vasoconstriction which combats the falling blood pressure elevation of the right ventricular pressure which helps to squeeze blood past the obstruction and elevation of the venous pressure which serves to encourage the right ventricle. It is as yet uncertain whether that chamber usually becomes overloaded or not. In cases which recover the embolus is gradually packed to the side of the vessels where it becomes organised and finally shrinks to a mere thread. Infarction of the lung does not necessarily occur because sufficient blood may pass through to nourish the tissues.

Subacute cases may occur in which repeated small emboli gradually block the pulmonary circulation over a period of weeks or months (Belt 1939).

Pulmonary infarction When an embolus lodges distally in a relatively small arterial trunk there is no rise of pressure in the pulmonary artery blood is not squeezed past the obstruction and the block is complete infarction of that part of the lung supplied by the occluded vessel follows (unless the collateral circulation is sufficient to nourish the ischaemic area). Of course such an event is likely to complicate massive pulmonary embolism and does so in 62 per cent of cases (Belt 1934) but it is a complication and not an essential part of the picture. Admittedly experimental pulmonary embolism does not cause infarction in animals unless the circulation is otherwise impaired (Karsner and Ash 1912) but no such condition appears to be necessary in clinical medicine. Infarcts of the lung are haemorrhagic because blood from the bronchial arteries exudes into the devitalised area. If this second source of nutrition is adequate for the needs of the tissue infarction does not occur. When the haemorrhagic zone reaches the surface of the lung a sero fibrinous pleural reaction develops pain may be severe as in any other pleurisy effusion is common and is usually blood stained.

Pulmonary infarcts are nearly always embolic in origin (Virchow 1856) very few are due to primary pulmonary thrombosis and they rarely complicate idiopathic pulmonary hypertension or Fallot tetralogy two diseases in which primary thrombosis is relatively common.

CLINICAL FEATURES

Massive pulmonary embolism In a typical dramatic attack the patient feels as if he had been struck in the centre of the chest and rapidly becomes faint grey cold clammy and breathless. Central sternal pain may be indis-

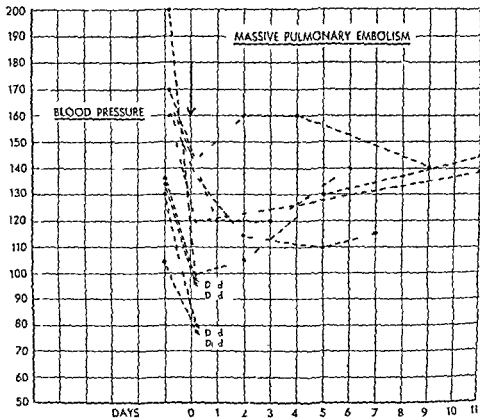


Fig 16 01—Behaviour of the blood pressure in 9 cases of massive pulmonary embolism. There is invariably a profound initial drop. In the group shown those with relatively high blood pressures previously recovered, whereas those with relatively low pressures previously died.

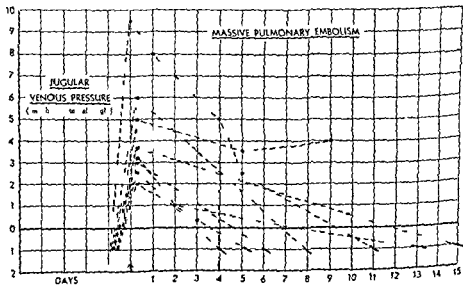


Fig 16 02—Behaviour of the venous pressure in 8 cases of massive pulmonary embolism. There is initial elevation in all, but it is rarely maintained for more than a few days.

tinguishable from that of acute myocardial infarction. Consciousness may be lost. Peripheral cyanosis is evident in the ears, lips and nail beds, but elsewhere pallor is usually more noticeable. Sweating is commonly profuse. The pulse is thready and rapid, or may be imperceptible; the blood pressure is low or immeasurable (fig. 16.01). The jugular venous pressure is invariably raised (fig. 16.02) and the liver may be palpable; cardiac oedema is not seen in acute cases, but may occur later in the subacute form. Examination of the lungs may reveal nothing abnormal. The heart sounds are usually soft, although the second sound at the base may be relatively accentuated or widely split (if there is right bundle branch block). Clinical and direct visual evidence of dilatation of the pulmonary artery proximal to the embolus have been described by McGinn and White (1933) and by Churchill (1934) respectively. The Graham Steele murmur of functional pulmonary incompetence has been heard (White and Brenner, 1933). Occasionally a pericardial friction rub develops over the base of the distended pulmonary artery (White, 1937).

Rarely patients die abruptly at the onset, presumably from reflex cardiac inhibition or ventricular fibrillation, such deaths being preventable by atropine in animals and being independent of the size of the embolus (Scherf and Schonbrunner, 1937). The great majority, however, survive the initial insult, but about one third die subsequently from circulatory obstruction, approximately 10 per cent within 10 minutes, 30 per cent within an hour, and 60 per cent in a matter of hours or days (de Takats and Fowler, 1945). On the other hand, about two thirds recover—within hours, days or weeks. Throughout this anxious period there is a 25 per cent risk of another, and perhaps fatal, embolus.

Massive pulmonary embolism, however, is not always dramatic, and mild cases are easily overlooked. Passing tightness of the chest, fleeting unexplained breathlessness, transient faintness, or a symptomless rise of systemic venous pressure may be the sole manifestation of an event that brought death very close.

Subacute cases may pass gradually into congestive heart failure without a single incident suggesting embolism; the clinical features of these rare cases resemble those of primary hypertensive pulmonary heart disease.

It should be noted that calling for the bed pan and falling back dead is not specially correlated with pulmonary embolism. The phenomenon appears to be associated with impending death from ventricular fibrillation or asystole, and may occur as a tragic climax to many forms of heart disease, including aortic stenosis and myocardial infarction. The colonic disturbance may be a vagal manifestation. Abrupt death from pulmonary embolism, preceded or not by a call to stool, is rare, as already mentioned.

The diagnosis of acute right ventricular stress may be proved electrocardiographically (fig. 16.03). Limb leads show sinus tachycardia, a constant S wave in lead 1, a frequent Q wave in lead 3, inversion of T_3 , flattening or slight inversion of T_2 , and rather low voltage (Barnes, 1937).

Occasionally P becomes tall and sharp (Wood 1948). The e appearances are not unlike those of posterior myocardial infarction although an absent S_1 , conspicuous Q_2 and elevation of the R-T segment in lead 3 should be sufficient to distinguish the latter in standard leads. Again Q_3 in cases of

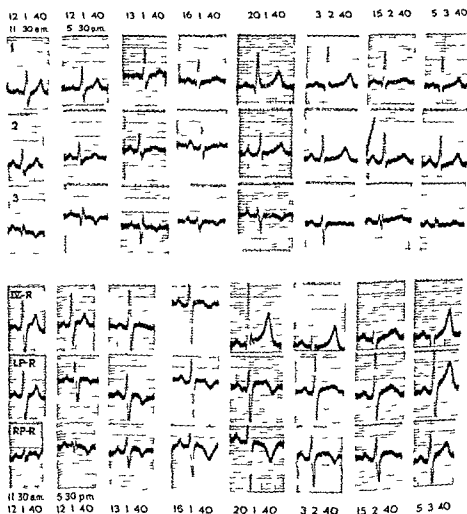


Fig. 16-03—Electrocardiogram showing the characteristic appearances associated with massive pulmonary embolism (lead IV-R=CR₄, LP-R=CR₂₋₃, RP-R=CR₁)

massive pulmonary embolism is caused by cardiac rotation and is not seen in lead VF. In multiple chest leads appearances are equally characteristic (Wood 1941) the T wave is nearly always inverted in leads V₁₋₃ over the right ventricle sometimes in V₄ and occasionally even in V₅ (fig. 16-03) and clockwise rotation or displacement of the interventricular septum to the left brings the RS pattern round as far as V₅ or even V₆. There are no pathological Q waves and the R-S-T segment is not deviated from the base

line but in about 15 per cent of cases there is transient right bundle branch block (fig 16 04). These changes are not immediate but develop within a few hours and are usually maximum within one to three days. Recovery is relatively slow, three to six weeks elapsing before the T wave is finally upright again in leads V₁ or V₂.

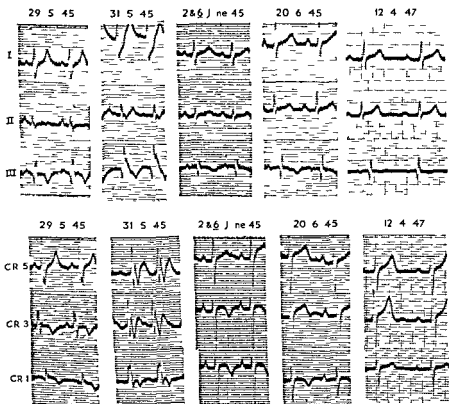


Fig 16 04—Electrocardiogram showing transient right bundle branch block in a case of massive pulmonary embolism

The electrocardiographic pattern has been variously attributed to reflex coronary spasm (Scherf and Boyd 1939) to interference with the coronary blood flow through the right ventricle owing to the combined effect of a low aortic and high right ventricular pressure (Durant Long and Oppenheimer 1947) and to right ventricular stress (Wood 1941). There is no proof of coronary spasm and although usually ascribed to a vagal reflex, the electrocardiographic pattern is unaltered by vagal section in dogs (Malinow Katz and Kondo 1946). In experimental air embolism in dogs Durant Long and Oppenheimer (1947) directly observed the development of right ventricular ischaemia. Their thesis is supported by the 60 per cent frequency of constricting substernal pain in clinical cases (Wood 1947).

22 7 38 4720



Fig. 16 05—Skiagram showing a small pulmonary infarct at the right base with a little hæmorrhagic effusion

On the other hand remarkably similar changes are found in any condition giving rise to right ventricular stress

Pulmonary infarction When an embolus lodges in a small or moderate sized pulmonary artery there are no immediate symptoms or signs. Within a variable time however, impossible to determine clinically a sudden pleural pain may signal the accident. Hæmoptysis may precede or follow the pain may not occur at all or may occur without pain. Coincident with these events and especially if there is pleurisy the respiratory rate rises and fever develops. Examination of the chest reveals little at first except perhaps pleural friction but within a day or two the percussion note may become impaired the breath sounds diminished and râles may be heard sometimes there are frank signs of consolidation or of fluid. When the diagnosis is in doubt a specimen of this fluid should be obtained for it is hæmorrhagic in most cases of infarction. A skiagram is also helpful the cone shaped infarct casting a triangular egg shaped or rounded shadow according to its lateral oblique or antero posterior lie (fig 16 03). Unfortunately the characteristic appearance is often obscured by the obliterating shadow of fluid. When embolism occurs without infarction skiagrams may show a segmental area of increased translucency owing to the absence of vascular shadows in the ischæmic area (Shapiro and Rigler 1948).

Pain hæmoptysis fever and tachypnœa may last a week or two but the patient does not look or feel seriously ill unless the primary disease makes him so. Moderate leucocytosis and a rapid erythrocyte sedimentation rate are the rule the former lasting a few days the latter several weeks as with cardiac infarction.

Whilst the above description applies to most cases it should be added that infarcts may be clinically silent and are often only discovered at necropsy. In other cases secondary infection complicates the picture or the embolus may be infected from the start pulmonary abscess or empyema may then develop.

Venous thrombosis In all cases of suspected pulmonary embolism a search should be made for the source. As previously stated this is commonly phlebothrombosis in the legs. It usually begins in the calf where there may be deep muscle tenderness or pain on dorsiflexing the foot (Homans' sign). Superficial thrombosis in the long saphenous vein may be felt as a solid cord and is usually tender. With thrombo phlebitis the overlying skin is hot red indurated and painful. Extension to the femoral vein causes a conspicuous rise of skin temperature in the affected limb a most useful sign of serious phlebothrombosis. œdema also occurs in many cases but is less constant.

PROGNOSIS

It is not easy to assess the true mortality rate in thrombo embolism for many mild cases are overlooked but in a series of twenty clinically recognised cases of massive pulmonary embolism seen by the author six died

In necropsy material about two thirds of all pulmonary emboli are major involving more than 50 per cent of the cross section of the pulmonary arteries (Belt 1939) but it is naturally the more severe ones that are seen at necropsy. From evidence of this kind it is estimated that nearly two thirds of all cases of massive pulmonary embolism recover and that less than a third of clinical thrombo emboli are massive this gives a total mortality rate of about 10 per cent.

TREATMENT

Prophylaxis is most important and should be directed towards accelerating the venous circulation in the legs and preventing the clotting process in bed ridden patients.

Breathing exercises, massage, frequent changes of position, active movements of the legs for specified times every day, prevention of dehydration and limitation of morphine are simple, logical and effective measures. Heart failure should be treated quickly and adequately. Rest in bed should never be prolonged unnecessarily.

Heparin is the quickest and safest anticoagulant but it is too expensive for routine prophylactic use. It should certainly be employed, however, as soon as the thrombo embolic state is recognised. For 23 per cent are multiple (Nygård *et al.* 1940-41) and not more than a quarter of cases of massive pulmonary embolism are fatal at the first insult (de Takats and Fowler 1945). Heparin may be given intravenously in doses of 50 mg (5 000 units) four to six-hourly by continuous intravenous drip in doses of 150 to 300 mg daily (50 to 100 mg to a pint of normal saline) or intramuscularly combined with 2 ml of 2 per cent procaine in doses of 150 mg twice daily. The last route is simple and effective. Procaine prevents pain and bruising is rarely serious. The dose of heparin should be regulated so that the clotting time is maintained at about two to three times the normal (Murray and Best 1938). Pitkin's menstruum (gelatin 18 per cent, dextrose 8 per cent, glacial acetic acid 0.5 per cent, distilled water to 100 per cent) as a vehicle for heparin is usually too painful for routine use.

Dicoumarol, the cause of hæmorrhagic sweet clover disease of cattle (Link 1943) is a cheaper and equally effective anticoagulant but its action is delayed for forty eight to seventy two hours and it is more difficult to control. It is given by mouth in a single dose each day, beginning with 300 mg on the first day, 200 mg on the second and 100 mg thereafter until the prothrombin activity of the blood approaches the desired level of 30 to 50 per cent of normal (de Takats and Fowler 1945). The dose is then reduced to 50 to 100 mg and is given daily or at less frequent intervals according to the prothrombin time. The latter must be measured daily during the first ten days of treatment then on alternate days for the effects of the drug cannot be accurately predicted and fatal hæmorrhage may occur following an overdose. Dicoumarol should be temporarily discontinued if there is hæmaturia, melæna or bleeding from other sites.

whatever the prothrombin time. Alarming hæmorrhage calls for fresh blood transfusion and repeated injections of 100 to 200 mg. of vitamin K intravenously (Barker *et al.* 1945).

In view of the delayed effect of dicoumarol, heparin is usually given as well during the first forty-eight to seventy-two hours. It is customary to continue dicoumarol for a month. With this treatment the post-operative mortality rate from massive pulmonary embolism in cases specially selected as thrombo-embolic risks has been reduced from perhaps 5 per cent to 0.1 to 1.0 per cent (Barker *et al.* 1945; Wright 1946).

Tromexan bis-3,3-(4-oxy-coumarinyl) ethyl acetate is likely to replace dicoumarol for it acts more quickly and is easier to control: the initial dose is 0.9 to 1.2 G. and the maintenance dose 0.3 to 0.6 G. daily (Burt, Wright and Kubik 1949).

Bilateral ligation of the femoral or common iliac veins or ligation of the inferior vena cava has been received with less enthusiasm, but it may be a life-saving procedure where anti-coagulants are contraindicated.

Treatment of acute pulmonary cardiopathy. Relatively mild cases recover spontaneously and require no special treatment. The majority of those clinically recognised, however, are seriously ill and require urgent attention. The objectives are to encourage the embolus to move on, or to be packed against the wall of the vessel, and to support the right ventricle. Eupavarine 1 mg. intravenously has been recommended, but it is doubtful whether it really dilates the main pulmonary artery or its larger branches. To encourage packing, the art is to play for time: if the patient can be kept alive (and every hour counts in his favour) progressive packing takes place automatically. To this end the vital centres must be supported, as the blood pressure is low; the patient should be nursed flat in order to encourage the cerebral circulation; oxygen, with the flow meter adjusted to six or seven litres per minute, should be administered continuously through a B.L.B. mask, or if possible the patient should be nursed in an oxygen tent, for the arterial oxygen saturation is often diminished; respiratory failure may temporarily respond to 0.48 G. of aminophylline, or to 10 ml. of coramine intravenously. Until recently it has been assumed that venesection and other venous pressure-lowering agents would help the right ventricle, but this is doubtful, for the raised venous pressure may be beneficial (fig. 16.06) (Wood 1947). Strophanthin 0.5 to 1.0 mg. intravenously helps to support the right ventricle.

The Trendelenburg operation—exposure of the pulmonary artery and removal of the clot—is only possible if a well-trained and thoroughly prepared surgical team is available, and is only practised when the situation is desperate: the operative mortality is over 90 per cent (Nygaard 1938) and spontaneous recovery is the rule rather than the exception. The first successful pulmonary embolectomy in Great Britain was reported by Ivor Lewis in 1939.

Treatment of pulmonary infarction. No specific treatment is required for

MASSIVE PULMONARY EMBOLISM

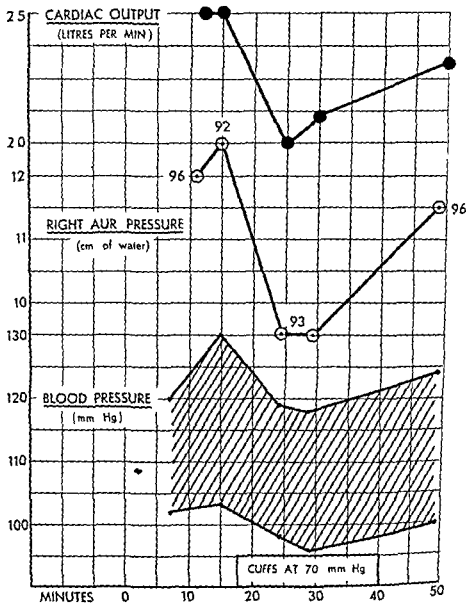


Fig 16 06—Effect of a venous pressure lowering agent (cuffs on the thighs) on the blood pressure and cardiac output of a case of massive pulmonary embolism

pulmonary infarction itself but secondary infection or septic embolism calls for sulphonamides or penicillin morphine may be necessary if there is severe pleural pain and hæmorrhagic pleural effusion may need aspirating if extensive Infarction does not contraindicate anticoagulants

PARADOXICAL EMBOLISM

Valvular patency of the foramen ovale is present in about a third of all individuals but the opening remains closed because the pressure in the left auricle is higher than that in the right When the right ventricle fails however the auricular pressures may be reversed the valve then opens and blood is shunted from right to left This event is improbable in heart failure secondary to mitral stenosis for the left auricular pressure remains too high Ideal conditions are presented by acute pulmonary heart failure due to massive pulmonary embolism for not only is the right auricular pressure then raised but the left is lowered as in pulmonary stenosis and emboli are already forthcoming Having passed through the foramen ovale the embolus is carried into the systemic circulation and may lodge in any cerebral visceral or peripheral artery

AIR EMBOLISM

Small quantities of air may be injected into the systemic venous system of healthy subjects with little risk indeed about 15 ml per kg body weight are required to kill a dog even when injected rapidly (Wolfe and Robertson 1935) Fatalities have occurred however when air has been accidentally introduced into a vein during an operation intravenous infusion therapeutic or diagnostic procedure The clinical features are those of massive pulmonary embolism but in addition a loud churning sound or millwheel murmur may be heard over the right ventricle and pulmonary artery Death appears to result from circulatory obstruction due to air lock in the outflow tract of the right ventricle Treatment consists of turning the patient prone and head down to bring the outflow tract below the level of the right ventricular cavity (Durant Long and Oppenheimer 1947) A similar manoeuvre has proved life saving in dogs but has not yet been tried in man

FAT EMBOLISM

Globules of fat may penetrate the systemic venous circulation following fractures usually of the femur and accidents have occasionally occurred during therapeutic or diagnostic procedures involving the use of oil Fat embolism has several characteristics which help to distinguish it from other forms First it happens within a few hours of the accident perhaps while manipulating the injured limb under anæsthesia or when moving the patient to the X ray department Second signs of multiple systemic embolism usually complicate the picture owing to the passage of fat globules

through the pulmonary capillaries. Thus there may be severe headache, drowsiness or loss of consciousness usually without localising signs. Multiple petechial spots may appear in the skin, red cells, albumin and droplets of oil may be found in the urine. Third, breathlessness and cyanosis are associated with the development of fine crepitations over all areas of the lungs, and skiagrams show an abundance of cotton wool shadows in all zones. The mortality rate is similar to that of other forms of massive pulmonary embolism, but those who survive recover remarkably quickly—often within forty-eight hours.

EMBOLISM DUE TO FOREIGN BODY

Metallic fragments from gun shot wounds and even bullets may enter the circulation in rare instances. Such an event should be considered if a skiagram shows an intra-thoracic foreign body when there is no wound of the chest or adjacent structures. An intravascular metallic foreign body may remain mobile for several days and may move against the bloodstream if so directed by the force of gravity. Surgical attempts to remove the missile may be foiled by such behaviour. An excellent example was described by Bauer (1943).

MALIGNANT EMBOLI

Cancer cells may infiltrate the systemic venous system and be swept into the lungs in the form of cellular emboli. Subacute pulmonary heart

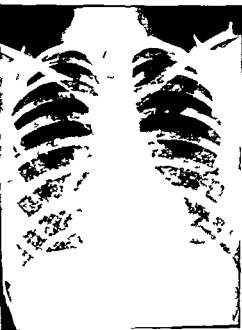


Fig. 16 07—Skiagram showing miliary embolic carcinomatosis of the lungs

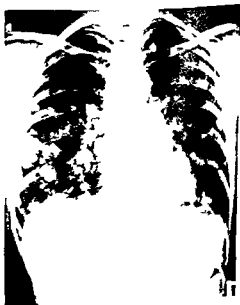


Fig. 16 08—Radiological appearances of the lungs showing embolic secondaries due to chorionepithelioma

(By courtesy of Dr. Philip Elliman)

disease develops if more than two thirds of the vessels are blocked the clinical features resembling those of massive pulmonary embolism but with an insidious onset and progressive course. The diagnosis may be suggested by the skiagram which may show minute miliary lesions (fig 16 07). Cases so far reported have been due either to carcinoma of the stomach (Brill and Robertson 1937) or breast (Mason 1940) or to chorion epithelioma (fig 16 08).

Subacute cor pulmonale may also be due to multiple pulmonary thromboses secondary to perivascular lymphatic carcinomatous infiltration (Brill and Robertson 1937).

REFERENCES

- Barker N W, Cromer H E, Hurn M and Waugh J M (1945) The use of dicoumarol in the prevention of post operative thrombosis and embolism with special reference to dosage and safe administration. *Surg* 17 207.
- Barnes A R (1937) Pulmonary embolism. *J Amer med Ass* 109 1347.
- Barne C G (1941) Electrocardiogram after pneumonectomy. *Proc Roy Soc Med* 34 606.
- Bauer H H (1943) Penetrating gunshot wound of the heart - triple embolism by a bullet. *Der chirurg* 15 697.
- Belt T H (1934) Thrombosis and pulmonary embolism. *Amer J Path* 10 129. — (1939) Late sequelæ of pulmonary embolism. *Lancet* ii 730.
- (1939) "The ætiology of lung infarction. *Brit Heart J* 1 283. — (1939) Autopsy incidence of pulmonary embolism. *Lancet* i 1259.
- Brill I C and Robertson T D (1937) Subacute cor pulmonale. *Arch intern Med* 60 1043.
- Burt C C, Wright H P and Kubik M (1949) Clinical tests of a new coumarin substance. *BMJ* ii 1250.
- Churchill E D (1934) The mechanism of death in massive pulmonary embolism. *Surg Gynec and Obstetr* 59 513.
- Collins D C (1936) Pulmonary embolism based upon study of 271 instances. *Amer J Surg* 33 210.
- Crutcher R R (1948) Venous thrombosis and pulmonary embolism. *Kentucky Med J* 46 427.
- Durant T M, Long J and Oppenheimer M J (1947) Pulmonary (venous) air embolism. *Amer Heart J* 33 269.
- Eppinger E C and Kennedy J A (1938) The cause of death in coronary thrombosis with special reference to pulmonary embolism. *Amer J med Sc* 195 104.
- Gibbon J H, Hopkinson M and Churchill E D (1937) Changes in circulation produced by gradual occlusion of pulmonary artery. *J clin Invest* 11 543.
- Hampton A O and Castleman B (1940) Post mortem chest teleroentgenograms with autopsy findings. *Amer J Roentgenol* 43 305.
- Homans J (1947) Venous thrombosis and pulmonary embolism. *New Engl J Med* 236 196.
- Karsner H T and Ash J E (1912-3) Studies in infarction II Experimental bland infarction of the lung. *J med Res* 27 205.
- Lewis I (1939) Trendelenburg's operation for pulmonary embolism. *Lancet* i 1037.

Lank K P (1943-4) The anticoagulant from spoiled sweet clover hay, Harvey Lectures p 162 Pennsylvania

McCartney J S (1945) Post operative pulmonary embolism *Surgery* 17 191

McGinn S and White P D (1935) Acute cor pulmonale resulting from pulmonary embolism its clinical recognition *J Amer med Ass* 104 1473

McPheeters H O and Rice C O (1928) Varicose veins complications direct and associated following injection treatment Review of the literature *Ibid* 91 1090

Macht D I (1946) Thromboplastic properties of some mercurial diuretics *Amer Heart J* 31 460

Malinow M R Katz I N and Kondo B (1946) Is there a vagal pulmonary coronary reflex in pulmonary embolism? *Ibid* 31 702

Mason D G (1940) Subacute cor pulmonale *Arch intern Med* 66 1221

Massie E Stillerman H S Wright C and Minnich V (1944) Effect of administration of digitalis on coagulability of human blood *Ibid* 74 172

Murray C D W and Best C H (1938) Use of heparin in thrombosis *Ann Surg* 108 163

Nygaard K K (1938) Consideration of clinical diagnosis and possibilities for the Trendelenburg operation *Proc Mayo Clin* 13 586 — (1941) Hemorrhagic diseases photoelectric study of blood coagulability London —

Barker N W Walters W and Priestly J T (1940) A statistical study of post operative venous thrombosis and pulmonary embolism I Incidence in various types of operation *Proc Mayo Clin* 15 769 — — — — —

(1941) A statistical study of post operative venous thrombosis and pulmonary embolism II Predisposing factors *Ibid* 16 1 — — — — — (1941)

A statistical study of post operative venous thrombosis and pulmonary embolism III Time of occurrence during post operative period *Ibid* 16 17

Pilcher R (1939) The role of obstruction in fatal pulmonary embolism *Lancet* 1 1257

Reitun F (1936) Thrombose und tödliche Lungenembolie *Lirchows Arch f path anat* 297 535

Scherf D and Boyd L J (1939) Cardiovascular diseases London —

and Schonbrunner E (1937) The pulmonary reflex in lung emboli *Klin Wchenschr* 16 340

Shapiro R and Rippler I (1948) Pulmonary embolism without infarction *Amer J Roentgenol* 60 460

de Takats F and Fowler E F (1945) The problem of thrombo embolism *Surg* 17 153

Virchow R (1856) Ueber die Verstopfung der Lungenarterie *Gesammelte als handlungen Frankfurt a m* 224

Westerborn A (1937) Über die Emboliegefahr bei Injektionsbehandlung von varizen nebst einem Bericht über die in Schweden vorkommenden Emboliefälle *Acta chir Scand* 321

White P D (1937) Heart disease New York — Brenner O (1933) Pathological and clinical aspects of the pulmonary circulation *New Engl J Med* 209 1261

Wolfe J B and Robertson H I (1935) Experimental air embolism *Ann intern Med* 9 162

Wood P H (1941) Pulmonary embolism diagnosis by chest lead electrocardiography *Brit Heart J* 9 21 — (1947) Discussion on pulmonary embolism *Ibid* 9 308 — (1948) Electrocardiographic appearances in acute and chronic pulmonary heart disease *Ibid* 10 87

Wright I S (1946) Practical considerations in the conservative treatment of thrombophlebitis *N Y J Med* 46 1819

CHAPTER XVII

PULMONARY HEART DISEASE

PULMONARY heart disease may be defined as a disorder of the heart resulting from disease of the lungs or of the pulmonary circulation. It may be acute as in massive pulmonary embolism, subacute as in recurrent thrombo embolism and secondary carcinomatosis of the lungs or chronic. The acute and subacute forms have already been discussed in the last chapter; the chronic form may be hypertensive or anoxic.

HYPERTENSIVE PULMONARY HEART DISEASE

Pathogenesis. Pulmonary hypertension associated with sclerosis of the pulmonary arteries is usually of unknown etiology (primary pulmonary vascular sclerosis or idiopathic pulmonary hypertension) but may sometimes be due to periarteritis (Fiskelund 1943). Bilharzia (Bedford *et al* 1946), recurrent pulmonary embolism (Castleman and Bland 1946) and less certainly to other known agents.

In idiopathic cases the arterial lesions reported in the literature include atherosclerosis, hypoplasia of the media (Gilmour and Evans 1946), great thickening of the intima (Barrett and Cole 1946) and widespread thromboses (Gold 1946); in others the pulmonary arterial tree has shown little structural abnormality (East 1940). Earlier reports reviewed by Brenner (1935) were similar. All agree that even when lesions are gross many vessels escape entirely and that advanced lesions may be found without hypertrophy of the right ventricle.

Various hypotheses have been offered to explain these findings: (1) the arterial lesions may be due to idiopathic pulmonary hypertension (East 1940); (2) congenital hypoplasia of the media may lead to protective intimal thickening which may initiate a vicious circle by causing pulmonary hypertension (Gilmour and Evans 1946); (3) both pulmonary hypertension and the arterial lesions described may result from repeated pulmonary embolism (Castleman and Bland 1946). It is probable that hypertensive pulmonary heart disease may be caused by several different agents and that the incidence of idiopathic cases will dwindle as more are recognised.

It may be noted here that the structure of the small pulmonary arteries and arterioles suggests a passive rather than an active role: thus there is no muscle in a pulmonary arteriole, merely a layer or two of elastic tissue around the endothelium. In the small arteries (0.1 to 1.0 mm. in external diameter) the muscular media averages only 14 per cent of the external diameter of the vessel compared with 36 per cent in a systemic artery of

the same size (Brenner 1935) The question therefore arises whether these vessels are capable of sufficient vasoconstriction to embarrass the right ventricle To this may be said that it is unsound to argue about physiological events in terms of anatomical structure Thus capillaries can constrict against astonishing pressures a function retained by pulmonary capillaries although they have no Rouget cells (Wearn, 1934) and there is no valid reason for supposing that small pulmonary arteries and arterioles are not possessed of considerable contractile power It is known that they are supplied with vasoconstrictor fibres through the sympathetic (Bradford and Dean 1894) there is evidence that they are supplied with vasodilator fibres also (Daly and Euler, 1932) and they may respond by constriction and dilatation to adrenergic and cholinergic drugs respectively (Young 1939) Direct evidence of pulmonary vasoconstriction in man has been obtained recently by Motley and Courmand (1947) who demonstrated a considerable rise in pulmonary arterial pressure during periods of artificial anoxia

Incidence The disease may occur in either sex and at any age including childhood but is rare out of 100 cases of chronic pulmonary heart disease analysed by the author only three could be classified as idiopathic

Physiology Pulmonary hypertension of any etiology imposes a mechanical burden on the right ventricle and pulmonary artery just as systemic hypertension affects the left ventricle and aorta the right ventricle hypertrophies and the pulmonary artery dilates

Sooner or later pure right ventricular failure develops the cardiac output is low and cyanosis is peripheral The arterial oxygen saturation remains normal until near the end but may then fall to about 80 per cent

Clinical features Idiopathic cases are rarely recognised until far advanced but evidence is accumulating which suggests the likelihood of a much longer course than generally believed

The findings are usually those of pure right ventricular failure with normal rhythm although auricular flutter or fibrillation may develop later The jugular venous pressure is raised and the a wave conspicuous there may well be functional tricuspid incompetence the liver is enlarged and tender and there is usually dependent oedema but the lungs are dry and there is no orthopnoea Cyanosis is often conspicuous but is associated with cold extremities a low cardiac output a normal or near normal arterial oxygen saturation and a high arterio venous oxygen difference it is therefore peripheral not central Polycythæmia and clubbing are usually absent

The cardiac impulse is diffuse and right ventricular in quality there may be *systolic lifting* in the third left intercostal space over the right ventricular outflow tract There is no loss of cardiac dullness rather the reverse and there are no other signs of emphysema There are usually no murmurs unless functional pulmonary incompetence develops but the pulmonary element of a normally split second heart sound is loud and high pitched and presystolic gallop may be heard to the left of the sternum

The electrocardiogram shows a pulmonary P wave and strong right ventricular preponderance (fig 17 01)

Fluoroscopy reveals a prominent pulmonary artery and right ventricle and though the proximal pulmonary vascular shadows may be heavy the more peripheral lung fields are clear (fig 17 02) Hilar pulsation may be seen but is unimpressive The right auricle is moderately dilated the left



Fig 17 01 (*left*)—Electrocardiogram showing prominent P waves and right ventricular dominance in a case of idiopathic pulmonary hypertension



Fig 17 0 —Skiagram showing prominence of the pulmonary arc and right ventricular enlargement in a case of idiopathic pulmonary hypertension

flat and there are no signs of emphysema Catheterisation reveals an extremely high pressure in the right ventricle and pulmonary artery (the mean pulmonary artery pressure ranged between 55 and 81 mm Hg in five cases of the author's) and no shunt

Diagnosis The features described make a characteristic and pathognomonic picture Pure pulmonary stenosis is excluded by the quality of the second sound and by the absence of a basal thrill atrial septal defect by lack of pulmonary plethora character of the second heart sound inconspicuous hilar pulsation and electrocardiogram anoxic pulmonary heart disease by the lack of emphysema obviously low cardiac output and peripheral rather than central cyanosis mitral stenosis by the absence of a mitral diastolic murmur and by the flat left auricle and different P wave

Diagnostic difficulty may arise however when pulmonary hypertension causes reversed interatrial shunt through a patent foramen oval as discussed on page 245

Prognosis The disease is always progressive and is usually fatal within two years of its recognition

Treatment The results of treatment are poor a low sodium diet mercurial digitalis, venesection and thiouracil may prolong life but the patient usually remains bed ridden and symptoms become increasingly difficult to control an oxygen tent does not help

ANOXIC PULMONARY HEART DISEASE

Definition The anoxic type of pulmonary heart disease invariably results from emphysema and depends upon the combination in varying degree of pulmonary hypertension and hypoxia with the emphasis on the latter

Pathogenesis The essential underlying pathology is emphysema however caused The great majority of cases are due to chronic bronchitis or bronchial asthma relatively few to bronchiectasis silicosis other forms of pneumoconiosis (Griggs Coggin and Evans 1939) pulmonary tuberculosis congenital cystic lung or kyphoscoliosis (Chapman Dilland Graybiel, 1939)

When the vital capacity becomes seriously reduced ventilation may become inadequate even at rest, the arterial oxygen saturation falls (McMichael and Sharpey Schafer 1944) and the carbon dioxide content of the arterial blood rises (Taquiñi Fasciolo Suarez and Chiodi 1947) Cyanosis is thus central in origin and breathlessness may be chemical rather than reflex The deficiency in gaseous exchange tends to be compensated by an increase in cardiac output (McMichael and Sharpey Schafer 1944) rather than by polycythæmia but there is also an increase in utilisation of available oxygen At the same time the pulmonary arterial pressure rises (Bloomfield *et al* 1946) probably reflexly as a result of pulmonary hypoxia (Motlev Cournand *et al* 1947) Secondary structural changes in the pulmonary arteries such as atheroma and intimal thickening develop sooner or later in most cases and may increase the hypertension

Incidence Anoxic pulmonary heart disease is much more common than statistical evidence at present indicates this is partly due to the casual attitude often adopted towards cases of chronic bronchitis and emphysema and partly to the fact that such cases are not usually sent to cardiovascular clinics the diagnosis is not easily made without special investigation unless there is heart failure and bronchopneumonia may obscure the cardiac factor terminally

The condition probably accounts for 5 to 10 per cent of all cases of organic heart disease It is at least five times more common in men than in women and about 75 per cent of the patients are over 50 years old (Spain and Handler 1946)

Clinical features The patient is usually a man neither very old nor very

young He commonly gives a history of bronchial asthma or of recurrent winter bronchitis for many years with increasing breathlessness over the last year or two and may have sought advice because of recent swelling of the legs Cross examination yields little further information he may have had attacks of tightness in the chest associated with breathlessness but not paroxysmal cardiac dyspnoea he may have had substernal discomfort but not true angina he may prefer to be propped up a little at night but usually raises no objection to lying flat

Physical signs Emphysema is usually obvious the chest is distended and moves little with respiration cardiac dullness is absent and the percussion note is generally tympanitic the breath sounds are faint Central cyanosis may be gross as in the case discussed by Ayerza or scarcely detectable It may be recognised in warm situations as in the conjunctivæ and inner sides of the lips where it is unlikely to be confused with peripheral cyanosis Polycythæmia and clubbing are rare The hands are warm capillary pulsation digital throbbing a modified water hammer pulse and increased pulse pressure may often be demonstrated Elevation of the jugular venous pressure and tachycardia may confirm the impression that the cardiac output is raised Papilloedema sometimes occurs (frontispiece)

The heart itself is apt to be camouflaged by over expanded lung the apex beat is impalpable the left cardiac border impossible to locate by percussion the heart sounds difficult to hear and the second sound at the base often inaudible there are no murmurs but right sided presystolic gallop may be heard or felt just to the left of the sternum in the fourth intercostal space

When there is true congestive failure the liver is distended and tender and dependent œdema is the rule

In severe cases vasomotor collapse is apt to occur when some superimposed broncho pulmonary infection lowers the arterial oxygen saturation relatively suddenly the blood pressure drops the pulse becomes small and thready the cardiac output low and the skin cold and clammy the outlook is then very grave

The electrocardiogram Emphysema alone does not materially affect the electrocardiogram although it may cause clockwise rotation about the antero posterior and longitudinal axes (viewed from the front and below) Thus there may be right axis deviation in standard leads an RS pattern in lead VL a QR pattern in lead VF and an RS pattern from V₁ as far as V₅ or even V₆ (fig 17 03) When the heart is exceptionally vertical VR and VL may be indistinguishable or backward tilting of the apex may cause VL to resemble an œsophageal lead from the back of the heart

In 100 cases of chronic pulmonary heart disease analysed by the author (Wood 1947) the following electrocardiographic appearances were found in *standard leads* (fig 17 04a to e)

Pulmonary P wave (fig 17 04)	85
Right axis deviation—	
with T ₃ (and often T ₂) inverted (fig 17 04a)	20
with T upright in all leads (fig 17 04b)	30

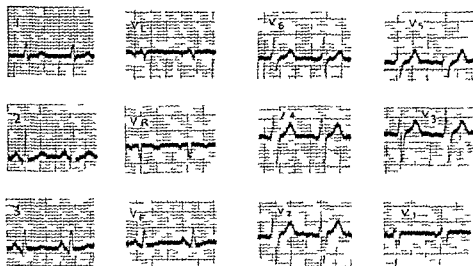


Fig 1703—Electrocardiogram in a case of emphysema showing a vertical electrical position and clockwise rotation (viewed from below)

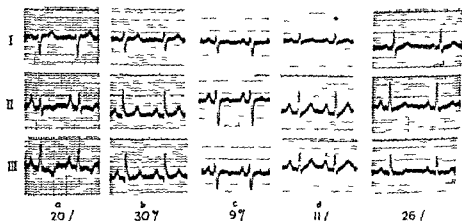


Fig 1704—Standard lead electrocardiographic findings in 100 cases of anoxic pulmonary heart disease

- (a) Right axis deviation with inversion of T₃ (and often T₂)
- (b) Right axis deviation with upright T waves
- (c) Dominant S wave in all standard leads
- (d) Tendency to right axis deviation
- (e) Normal QRS axis

The pulmonary P wave is seen in all

Prominent S wave in all leads (fig 17 04c)	9
Tendency to right axis deviation (fig 17 04d)	11
Normal axis of QRS (fig 17 04e)	26
Right bundle branch block	4
Low voltage	40

Multiple chest leads revealed the following

Normal QRS deflections in the majority (fig 17 05a and b)

	Per cent
Inversion of T from V ₁ -V ₃ (fig 17 05c)	13
Dominant R wave in V ₁ with conspicuous S in V ₅ (fig 17 05d)	16
Dominant S wave from V ₁ -V ₅ (fig 17 05e)	16

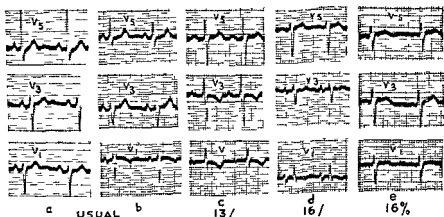


Fig 17 05—Chest lead findings in anoxic pulmonary heart disease

- (a) (b) Normal chest leads
(c) Inversion of T from V₁ to V₃
(d) Dominant R wave in V₁ with conspicuous S in V₅
(e) Dominant S wave from V₁ to V₅

Unipolar limb leads nearly always showed a vertical electrical position

Particular attention is drawn to the frequency and importance of the pulmonary P wave (fig 17 06 and 17 07). In normal controls the maximum auricular deflection very rarely measures more than 1.5 mm in amplitude and averages 1 mm (fig 17 08). The pulmonary P wave commonly ranges between 2 and 3 mm in height but is never widened. It is not seen in normal vertical hearts which refutes the suggestion that it depends on cardiac rotation due to emphysema. It cannot be attributed to anoxia for it is an early finding and tends to diminish in voltage when anoxia becomes severe nor is it present in cases of severe anaemia. It cannot be ascribed to an elevated cardiac output for it is seen in a much less conspicuous form in cases of thyrotoxicosis in which the cardiac output is considerably higher moreover as already mentioned P is of low voltage in severe

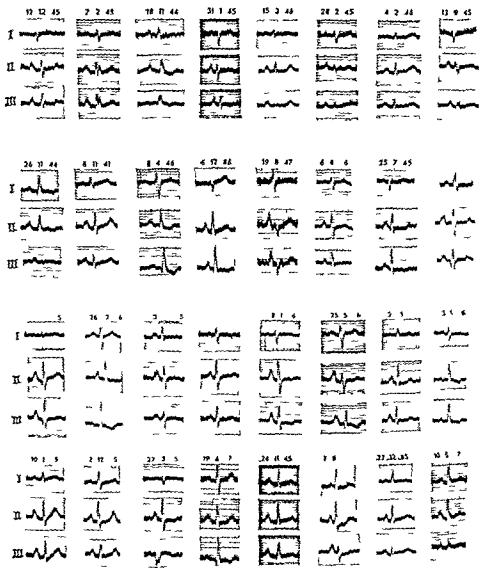


Fig 17 06—Standard limb lead electrocardiograms of 32 unselected cases of anoxic pulmonary heart disease with relatively low voltage showing the frequency amplitude and shape of the pulmonary wave

chronic anaemia with outputs up to 14 litres per minute. Intracardiac pressure studies have revealed little correlation between this P wave and the right auricular pressure but there appears to be some association be-

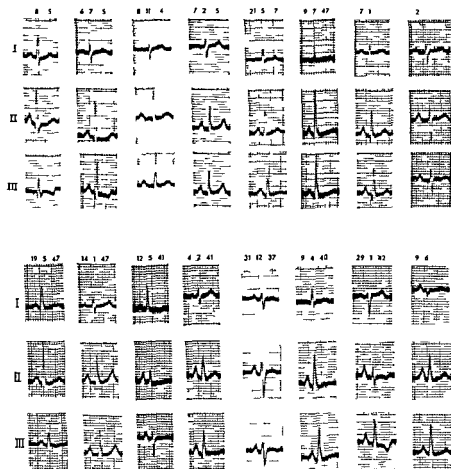


Fig. 17.07—Standard limb lead electrocardiograms of a further 16 unselected cases of anoxic pulmonary heart disease with normal voltage showing the frequency, amplitude and shape of the pulmonary P wave.

tween it and the right ventricular pressure. Just on what such a relationship may depend is unknown.

The pulmonary P wave is probably the earliest sign of cardiovascular disturbance resulting from emphysema or at least competes in this respect with elevation of the right ventricular pressure and slight reduction of the arterial oxygen saturation. It may develop several years before the onset of heart failure.

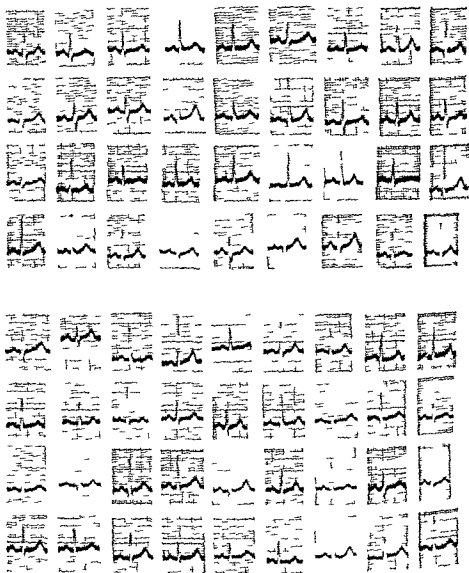


FIG. 17-08.—The maximum P waves in one or other of the standard leads (usually, lead 2) of unselected normal controls are shown for comparison with FIGS. 17-06 and 17-07.



(a)



(b)

Fig 17 09 (a) (b)—Skiagrams of two advanced cases of anoxic pulmonary heart disease showing dilatation of the pulmonary arc and of the left and right branches



(a)



(b)

Fig 17 10 (a)—Right anterior oblique position showing the increased density and diameter of the pulmonary artery at its bifurcation
(b) Left anterior oblique position showing the left pulmonary artery forming an almost straight line and as large as the aortic arch

Fluoroscopy Prominence of the main branches of the pulmonary artery at the hila with or without dilatation of the main pulmonary arc is seen in over 50 per cent of cases of severe emphysema (Parkinson and Hoyle 1937) but the changes are rarely conspicuous until pulmonary heart disease is well advanced (fig 17 09 and 17 10) Associated hypertrophy of the right ventricle is less easily demonstrated

Pulsation of the pulmonary artery and its main branches may be seen sometimes but does not compare with that in atrial septal defect and as a rule is absent Peripheral vascular markings are relatively unimpressive Enlargement of the right auricle is rare in the absence of failure The left auricle is flat and a prominence on the left border of the heart between the pulmonary and left ventricular arcs is never seen Owing to the raised cardiac output and to the frequency of coincident essential hypertension the aortic knuckle is usually well seen and may be unduly prominent

Finally there may be evidence of emphysema widening of the rib spaces elevation of the ribs and clavicle depression of the diaphragm and increased translucency of the lung parenchyma However emphysema is not cor pulmonale and too much should not be deduced from its presence

Special investigations The vital capacity is greatly reduced and is usually below 1500 ml The residual air is increased proportionately the total lung volume remaining normal Central cyanosis may be proved by arterial puncture the arterial oxygen saturation is usually between 60 and 80 per cent but may be much lower

The pulmonary artery and right ventricular pressures may be measured by means of cardiac catheterisation and are usually moderately raised but less so than in primary pulmonary hypertension

In 19 cases of emphysema without right ventricular failure Courmand and his colleagues found the systolic right ventricular pressure was normal (18 to 30 mm Hg) in 5 and between 34.5 and 57.5 mm Hg in 14 (Bloomfield *et al* 1946) They regard this rise as the earliest evidence of pulmonary heart disease

Cardiac outputs commonly range between 5 and 9 litres per minute (McMichael and Sharpev Schafer 1944) and do not seem to reach the high levels encountered in anaemia thyrotoxicosis and large arterio venous shunts

The right auricular pressure tends to be raised when the cardiac output is increased and may be high in clinical congestive failure On the other hand in some cases of emphysema it is remarkably low the mean pressure being less than minus 10 cm of saline below the sternal angle this may be attributed to anterior displacement of the sternal angle and to an unusually low intrathoracic pressure

Diagnosis The usual clinical problem is to decide whether the cardiovascular system is involved in a known case of emphysema but difficulty may also arise in distinguishing pulmonary heart disease from other cardio pathies and especially in unravelling a mixed etiology Other hyperkinetic

circulatory states may have to be excluded particularly beri beri in alcoholics but also thyrotoxicosis secondary carcinomatosis of the liver and Paget's disease of bone in emphysematous subjects. The commonest mixed etiology is the association of emphysema and hypertension. Although the differences between hypertensive heart failure and pulmonary heart failure are many it should be remembered that both may occur at the same time.

In the stage of low blood pressure and reduced cardiac output clinical diagnosis may be even more difficult. Toxic vasomotor collapse from bronchopneumonia may cause confusion, mitral stenosis, Pick's disease, atrial septal defect, mediastinal tumour, massive pulmonary embolism and many other conditions may have to be considered. The correct diagnosis can usually be made after full investigation but the first clinical impression can be very misleading.

Prognosis The diagnosis of chronic anoxic pulmonary heart disease usually carries with it a grave prognosis, few cases surviving a year but such diagnoses are rarely made before the onset of failure. With the newer methods of investigation circulatory involvement should be recognised much earlier perhaps by 5 or 10 years and appropriate treatment might then prolong life.

Treatment Vigorous preventive and symptomatic treatment of bronchitis and asthma may delay the development of serious emphysema indefinitely. Half-hearted measures must be condemned when the ultimate fate of these patients is realised.

By the time the cardiovascular system is involved emphysema is usually far advanced. A partly reversible state may be encountered however when acute bronchitis, bronchopneumonia or an asthmatic bout is superimposed on chronic changes of only moderate degree. In such cases infection should be treated promptly with penicillin or other forms of chemotherapy and bronchial spasm relieved by a dust free atmosphere and antispasmodics.

Although details of such treatment cannot be considered in a work of this kind one or two observations are necessary. Morphine is frequently lethal owing to its depressing effect on respiration, pethidine may quieten a restless patient just as well, is a good antispasmodic and does not depress respiration. Subcutaneous adrenaline is still the most effective way of relieving bronchial spasm, newer remedies such as the antihistamine drugs may be given in addition but not as a substitute. Isopropyl *nor* adrenaline which may be administered in sublingual tablets in doses of 20 to 40 mg is a useful preparation. Antispasmodics that improve the cardiac output or coronary circulation such as aminophylline may be chosen in preference to those that do not.

Whether the case is complicated by infection and bronchial spasm or not it is vitally important that the patient should be nursed in an oxygen tent. The effect of improving the arterial oxygen saturation is often dramatic, it prevents fatal vasomotor collapse, reduces the work of the heart and may lower the pulmonary blood pressure.

Mersalyl a low sodium diet and venesection should be used with caution. Howarth, McMichael and Sharpey Schafer (1947) have shown that in most cases with raised cardiac outputs the venous pressure is already at an optimum level and that lowering it by any means may reduce the output and harm the patient. In a minority, however, the heart is overloaded and then responds to such treatment in the usual way. Although clinically it may not be easy to judge the physiological state of the circulation, warm extremities and a full bounding pulse contraindicate all venous pressure lowering agents, whereas cold extremities, a small pulse and low blood pressure demand them (when the venous pressure is raised). When œdema is gross and the jugular venous pressure over 7 cm. saline above the sternal angle, mersalyl and a low sodium diet should probably be tried. Venesection is best avoided in all cases, as it is too drastic and may prove fatal if ill judged. Digitalis or strophanthin may be used without fear when the usual clinical indications are present.

If, after relief of bronchial spasm and infection, the vital capacity remains critically low and the arterial oxygen is below 80 per cent when the patient is out of the tent, thiouracil should be seriously considered as a means of reducing the oxygen requirement (page 385).

OTHER FORMS OF PULMONARY HEART DISEASE

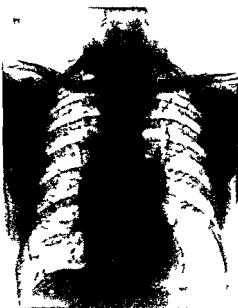
Ayerza's disease. Much confusion has arisen from the use of this term; it has been applied to cases of intense cyanosis and polycythæmia associated with syphilitic or other disease of the pulmonary arteries (Boyd, 1931). The facts are that Ayerza of Buenos Aires, in an unpublished clinical lecture (1901) described a single case of heart failure in which the patient was so cyanosed as to be almost black—a cardiac negro. Autopsy revealed much enlargement of the right side of the heart, dilatation of the bronchi and peribronchitis. Neither syphilis nor the state of the pulmonary vessels was mentioned. Arriaga (1913, 1924) was perhaps chiefly responsible for stressing the syphilitic origin of such cases, although other authors from the Argentine believed the arterial lesions to be atherosclerotic. Brenner (1935), after reviewing the evidence, concluded that there was no good reason for retaining the term Ayerza's disease on the grounds that published cases described nothing but chronic cor pulmonale.

Pulmonary heart disease associated with deformities of the chest. Gross kyphoscoliosis accounts for perhaps 5 per cent of cases of chronic cor pulmonale. The condition is associated with extensive collapse atrophy of part of the lung and severe emphysema of the remainder. Cardiovascular involvement is similar in type to that associated with other forms of pulmonary disease complicated by emphysema; kinking of the aorta (Corvisart) plays no part in its development.

The average age of death in these cases is about 50 years. A curious form of syncope has been described in a number (Chapman, Dill and Graybiel

1939) possibly due to sudden lowering of the right auricular pressure consequent upon compression of the inferior vena cava in certain postures (page 196)

Aneurysm of the pulmonary artery Aneurysm of the pulmonary artery is rare being found in less than 01 per cent of all autopsies and accounting for less than 05 per cent of all aneurysms (Deterling and Clagett 1947) The sexes are represented equally and about one third of the patients are



(a) 2nd March 1944



(b) 5th December 1946

Fig 17 11—Development of aneurysmal dilatation of the right pulmonary artery in a case of anoxic cor pulmonale

under 30 years of age (Boyd and McGarack 1939) The etiology is believed to be a congenital defect in the wall of the pulmonary artery in about 40 per cent syphilis in 30 per cent and chronic cor pulmonale with atherosclerotic pulmonary arteries in 30 per cent The diagnosis may be obvious on fluoroscopy if gross pulsation is seen if not it may be proved by means of angiocardiology (Robb and Steinberg 1940)

In pulmonary heart disease aneurysmal dilatation may develop remarkably quickly (fig 17 11) underlying congenital weakness of the arterial wall is difficult to exclude Thrombosis may occur in the sac or the whole vessel may be occluded but apart from such a complication the aneurysm is unlikely to influence the course of the primary disease Rupture is very rare

REFERENCES

- Arrillaga F C (1913) Sclerose de l'artere pulmonaire secondaire a certains etats pulmonaires chroniques (cardiaques noirs) *Arch d mal du Cœur* 6 518
- (1924) Sclerose de l'artere pulmonaire (cardiaques noirs) *Bull et mem Soc med d hop de Paris* 1 29.
- Barrett A M and Cole L (1946) Pulmonary vascular sclerosis with right ventricular failure *Brit Heart J* 8 76
- Bedford D L Arraras S M and Giris B (1946) Bilharzial heart disease in Egypt Cor pulmonale due to Bilharzial pulmonary endarteritis *Ibid* 8 87
- Bloomfield R A Lauson H D Cournand A Breed E S and Richards D W (1946) Recording of right heart pressures in normal subjects and in patients with chronic pulmonary disease and various types of cardio circulatory disease *J clin Invest* 25 639
- Boyd L J and McGavack T H (1939) Aneurysm of the pulmonary artery a review of the literature and a report of two cases *Amer Heart J* 18 562
- Boyd W (1931) The pathology of internal diseases Philadelphia
- Bradford J R and Dean H P (1894) The pulmonary circulation *J Physiol* 16 34
- Brenner O (1935) Pathology of the vessels of the pulmonary circulation part 2 part 5 *Arch intern Med* 56 1189
- Castleman B and Bland F F (1946) Organised emboli of the tertiary pulmonary arteries An unusual cause of cor pulmonale *Arch Path* 42 581
- Chapman E M Dill D B and Graybiel A (1939) The decrease in functional capacity of the lungs and heart resulting from deformities of the chest pulmonocardiac failure *Medicine* 18 167
- Daly I de B and Euler V von (1932) Functional activity of vaso motor nerves to lungs in dog *Proc Roy Soc Med* 110 9.
- Deterling R A and Clagett O T (1947) Aneurysm of the pulmonary artery review of the literature and report of a case *Amer Heart J* 34 471
- East T (1940) Pulmonary Hypertension *Brit Heart J* 2 189
- Eskelund V (1943) Periarteritis nodosa der pulmonalarterie und primäre pulmonalsklerose *Acta path et microbiol Scandinav* 19 13
- Gilmour J R and Evans W (1946) Primary pulmonary hypertension *J Path Bact* 58 687
- Gold M M A (1946) Congenital dilatation of the pulmonary arterial tree *Arch intern Med* 78 197
- Griggs D F Coggin C B and Evans N (1939) Right ventricular hypertrophy and congestive failure in chronic pulmonary disease *Amer Heart J* 17 681
- Howarth S McMichael J and Sharpey Schafer E P (1947) Effects of oxygen venesection and digitalis in chronic heart failure from disease of the lungs *Clin Sc* 6 187
- McMichael J and Sharpey Schafer E P (1944) The action of intravenous digoxin in man *Quart J Med* 13 123
- Motley H L Cournand A Werko L Himmelstein A and Dresdale D (1947) The influence of short periods of induced acute anoxia upon pulmonary artery pressure in man *Amer J Physiol* 150 315
- Parkinson J and Hovle C (1937) The heart in emphysema *Quart J Med* 6 59
- Robb G P and Steinberg I (1940) Visualisation of the chambers of the heart the pulmonary circulation and the great blood vessels in man summary of method and results *J Amer med Ass* 114 474

Spain D M and Handler B J (1946) Chronic cor pulmonale—sixty cases studied at necropsy *Arch intern Med* 77 37

Taquini A C Fasciolo J C Suarez J R E and Chiodi H (1947) Circulatory adaptations in Ayerza's syndrome—black cardias *Amer Heart J* 34 50

Wearn J T (1934) Normal behaviour of pulmonary blood vessels with observations on intermittence of flow of blood in arterioles and capillaries *Amer J Physiol* 109 236

Wood P H (1947) Electrocardiographic appearances in acute and chronic pulmonary heart disease *Brit Heart J* 10 87

Young R A (1939) The pulmonary circulation—before and after Harvey
The Harveian Oration London

THYROTOXICOSIS AND THE HEART IN MYXCEDEMA

THYROTOXIC HEART DISEASE

THE cardiovascular system is clearly involved from the onset of thyrotoxicosis although the term thyrotoxic heart disease is usually reserved for the late stage when auricular fibrillation or congestive heart failure dominates the scene. Such a distinction is artificial and simply means that a young and healthy heart can maintain a high output for years without distress but that an aged heart cannot.

Historical note Thyrotoxic heart disease was first adequately described by Caleb Hillier Parry (1815-1825) of Bath, who witnessed his first case in 1786. Flajani's publication of the details of one case (1802) appeared first but cannot be compared with Parry's account. Graves' description (1835) is also inferior. Carl von Basedow (1840) a general practitioner at Merseburg, Germany, called special attention to exophthalmos and drew a vivid picture of most of the features of primary exophthalmic goitre as we see it today, omitting only tremor which was later recognised and added to the Merseburg triad (exophthalmos, goitre and palpitations) by Pierre Marie (1883). For further historical details the reader is referred to the classical monographs of Cecil Joll (1932) and of Means and Richardson (1938).

NATURE OF THYROID HORMONE

The exact composition of thyroid hormone is not yet known. In 1893 Baumann obtained from thyroid tissue a protein-free physiologically active substance containing 10 per cent of iodine which he called iodothyrim. In 1899 Oswald showed that the active principle stored in the gland was attached to a protein in the form of thyroglobulin, this is the chief constituent of colloid. Kendall isolated thyroxine in 1915, showed that it contained 65 per cent of iodine, and demonstrated its potency. These researches culminated in the synthesis of thyroxine by Harington and Barger in 1927.

Thyroxine, however, accounts for only 40 to 50 per cent of the total iodine in the thyroid gland, is relatively insoluble and is not believed to be identical with thyroid hormone. The rest of the thyroid iodine is found in the practically inert substance diiodotyrosine, a likely precursor of thyroxine. According to Harington (1933) thyroxine and diiodotyrosine are probably linked with amino acids as constituents of thyroglobulin in colloid and the natural thyroid hormone is perhaps a thyroxine—containing peptide.

PATHOLOGY OF GOITRE

The normal thyroid gland consists essentially of numerous acini lined with epithelium and containing colloid material rich in iodine from which thyroid hormone appears to be liberated according to the demand. When the gland is stimulated the epithelium assumes an active columnar form and colloid tends to disappear. When there is little or none left the walls of the acini may become crenated like any other vesicle whose contents have been removed. In this phase the gland as a whole is soft and vascular and is not enlarged. When the stimulus ceases involution takes place the epithelium flattens, colloid reappears and the acini become distended. This is the resting phase and is characterised by a firmer, less vascular gland of somewhat larger size. If the stimulus to activity is excessive the morphological changes described above are supplemented by true hyperplasia of the acinar epithelium and subsequent involution may be incomplete leading to permanent enlargement of the gland.

Simple goitre is due to benign hyperplasia and develops when iodine supplies are short or diverted especially when thyroid demands are heavy (Marine 1927). This response to iodine lack is believed by some to be mediated by the production of excessive amounts of thyrotropic hormone from the anterior pituitary. Endemic goitre due to lack of iodine in the soil occurs in New Zealand, parts of Italy and North America and in many other mountainous districts or places remote from the sea. Iodine diversion may be due to polluted water (Marine and Enhart 1910, McCarrison 1927). Increased demands for thyroid hormone occur at puberty and during pregnancy.

Colloid goitre represents the resting involuted phase of previous benign hyperplasia (Marine 1930). When the stimulus subsides colloid reaccumulates in the acini, intervening walls between distended crowded vesicles break down to form cysts and the whole gland becomes tense and big. This process is innocent and causes no symptoms except possible discomfort in the neck.

In primary Graves disease persistent uncontrolled stimulation of the thyroid gland of unknown cause leads to marked hyperplasia and to wild manufacture and liberation of excessive amounts of thyroid hormone. The acinar epithelium is columnar and proliferated, the walls of the acini markedly crenated and the colloid practically all gone. The gland as a whole is soft, vascular and enlarged.

Nodular goitre is usually regarded as the end result of repeated cycles of hyperplasia and incomplete involution. The process probably begins with failure of complete involution of a previously stimulated and hyperplastic gland. Subsequent stimulation leads to local hyperplasia of these hypo-involuted nests and subsequent involution to local nodules of colloid goitre. Such a process may be repeated indefinitely. Thyrotoxicosis from nodular goitre depends chiefly upon the activity of the hyperplastic nests, the nodules themselves being mostly inert. The term adenomatous goitre

is therefore incorrect when applied to this type of lesion, and should be reserved to describe those cases in which thyroid nodules (usually single) are composed of solid masses of cells of fetal type. Compared with primary Graves disease nodular goitre usually runs a longer and less dramatic course, which by its very nature is necessarily phasic periods of activity alternating with periods of relative quiescence. Why production of thyroid hormone should exceed the demand is no more understood than it is in primary Graves disease. The implication of the anterior pituitary thyrotropic hormone may explain part of the mechanism but in no way solves the problem.

Physiology of the circulation under the influence of thyroxin The administration of thyroxin to man and mammals is followed after a time lag of several days by an appreciable rise in the basal metabolic rate. The increased oxygen requirement is met by elevation of the cardiac output not by greater utilisation of available oxygen (as occurs when the B.M.R. is raised by dinitrocresol) nor by polycythæmia. The high minute output is maintained more by tachycardia than by a raised venous pressure the stroke volume being but little increased (Friedberg and Solval, 1937). The strength of cardiac contraction is probably enhanced. These effects are usually attributed to the direct action of thyroxine on the heart.

At the same time the peripheral blood flow is greatly increased there is obvious vasodilatation in the skin, and adrenergic responses are magnified.

Morbid anatomy of the thyrotoxic heart There are no macroscopic changes in the thyrotoxic heart prior to the onset of auricular fibrillation and failure until then the heart weight remains normal. Cases exhibiting cardiac embarrassment during life may still show little at necropsy except some increase in heart weight and evidence of congestive failure (Kepler and Barnes 1932). In a few however there are scattered foci of fibrosis (Rake and McEldacharn 1932).

CLINICAL FEATURES

The hyperkinetic circulation of primary Graves disease is usually well tolerated because the subjects are young but in middle aged or elderly people with toxic nodular goitre cardiac embarrassment is the rule. The sex ratio favours women in the proportion of about 6:1 (Fraser and Dunhill 1934). A family history of goitre is found in 45 per cent of cases (Bruun 1945). Contributory factors include pregnancy the climacteric infection (such as tonsillitis) and perhaps emotional shock although the scarcity of thyrotoxicosis amongst active service casualties in the first two world wars was noteworthy. The role of iodine has already been discussed.

Of the symptoms loss of weight, heatintolerance, agitation or restlessness, palpitations and fatigue are the most important. Loss of weight as associated with a voracious appetite is particularly suggestive. Palpitations may be due to vigorous and rapid action of the heart or to paroxysmal auricular fibrillation the latter is especially significant.

Whilst the symptoms themselves are important the manner in which they are told and the general behaviour and appearance of the patient are often more so. The subject is usually a woman: she is commonly thin and talks quickly, often gesticulating to lend emphasis to her remarks. She may wear a scarf to hide an unsightly swelling in her neck, but her clothing is otherwise light. One of Parry's patients liked to sit in a draught, stripped to the waist, in order to keep cool (Parry, 1815). A good moment to look for the goitre is towards the beginning of the interview, when the patient may lean forward in her chair and swallow once or twice in nervousness. The eyes are characteristic, not so much because of exophthalmos, which is usually absent, but because of their typical stare. The trend of the patient's conversation is often illuminating and in sharp contrast to that of the anxiety neurotic. The latter complains of symptom after symptom in a challenging fashion, exaggerating their severity and stressing his inability to cope with them. The thyrotoxic patient tries to explain away her symptoms: she feels the heat, but of course it has been very warm recently; she is losing weight, but she supposes she was too fat before; she gets tired and irritable, but she knows she tries to do too much, and so on.

Physical examination may reveal a wealth of signs which are all directly or indirectly attributable to excess of thyroid hormone, except exophthalmos and goitre. They may be suitably described under four main headings:

1. *The eyes.* Exophthalmos may be present (fig. 18.01) but is uncommon in toxic nodular goitre. It is occasionally unilateral (fig. 18.02). Artificial glass eyes may also become proptosed. Its mechanism is still a subject of controversy (Zondek and Ticho, 1945) but exophthalmos is certainly not due to sympathetic stimulation, for it is not relieved by sympathectomy (Shaw, 1949) nor is it due to excess of thyroid hormone, which never reproduces it. Moreover, exophthalmos occasionally becomes more marked after thyroidectomy or treatment with thiouracil. In severe cases of exophthalmic ophthalmoplegia and malignant exophthalmos, thyrotoxicosis may be minimal, and the protrusion of the eye ball appears to be secondary to intense œdema of the orbital contents (Brain and Turnbull, 1938). Of great interest is the exophthalmos that can be produced in guinea pigs (also in rabbits and fish, but not so far in man) by injecting thyrotropic hormone, especially if the thyroid gland is first removed (Marine and Rosen, 1933). All these facts point to the likelihood of the pituitary being directly responsible, and provide further evidence that thyrotoxicosis may depend upon a primary pituitary disorder.

Retraction of the upper lid (fig. 18.03), revealing the white sclerotic above the iris (Dalrymple's sign), which may be unilateral, is also uncommon in toxic nodular goitre. It should be distinguished from exophthalmos, which reveals the white sclerotic below the iris by mechanically displacing the lower lid (Pochin, 1937-8).

If the patient looks up and then lowers the eyes to watch a descending object, the upper lid lags behind the movement of the eye ball, revealing



(a)

Fig. 18 01—Exophthalmic goitre
The first photograph (a) (in gipsy
dress) was taken in 1933 the second
(b) in 1936 The white sclerotics are
seen below the iris due to mechanical
displacement of the lower lid



(b)



FIG. 1802.—Unilateral lid retraction and exophthalmos.



Fig 18 03--Lid retraction and characteristic thyrotoxic stare

the white sclerotic above the iris (von Graefe 1864) Lid lag and lid retraction were for a long time attributed to stimulation of the sympathetic reinforcement of the levator palpebræ superioris (von Graefe 1864) but if sympathetic stimulation were responsible the lower lid would also be retracted which it is not (Pochin 1937-8 1939) Moreover both exophthalmos and lid retraction may occur when the ocular sympathetic is paralysed (Brain 1939) In the light of these findings von Graefe's hypothesis is untenable

The characteristic stare has already been mentioned It is more than lid retraction and infrequent blinking (Stellwag sign) it is a look which may occur independently and which can be recognised with experience The other eye signs of the textbooks are less important failure to wrinkle the forehead when the eyes are cast up (Joffroy's sign) may depend upon lid retraction and exophthalmos divergent strabismus as the eyes focus on an approaching object (Moebius sign) may be due to weakness of the oculomotor muscles as a result of stretching

2 *The hands* The hands are warm pink and slightly moist on both surfaces they are restless and expressive and may show a fine even constant tremor In contrast the hands of a psychoneurotic are cold and clammy being wet on the palms but not at the back they tend to be inert and expressionless tremor is coarse irregular and inconstant



Fig 18 04—Substernal goitre revealed by X rays

3 *The goitre* If a goitre is not seen it may be discovered by palpation It is best to stand behind the patient and to place the thumbs behind the sternomastoids and the fingers in front On asking the patient to swallow a nodular swelling may be felt moving upwards Posterior enlargement may be detected readily with this technique Practically all cases of thyrotoxicosis have a goitre although it is sometimes difficult to demonstrate (so called

masked hyperthyroidism) In such instances it may become more convincing after a course of Lugol's iodine Occasionally it is substernal and may be revealed by fluoroscopy (fig 18 04)

The goitre of thyrotoxic heart disease is commonly nodular irregular and asymmetrical It may displace the trachea to one side and the common carotid artery to the other and on rare occasions it may compress the trachea causing cough dyspnoea and stridor Sudden enlargement is usually due to hæmorrhage within a nodule or cyst Degenerated nodules may become calcified

Primary exophthalmic goitres are uniformly enlarged smooth and fleshy

They are similar to simple hyperplastic goitres, but more vascular. Sometimes an arteriovenous continuous thrill and murmur may be detected over the gland. Colloid goitres are also smooth and symmetrical but they are harder and as a rule larger. After a course of iodine primary exophthalmic goitre may feel like colloid goitre. Nodular goitre should be distinguished from other causes of thyroid enlargement and from other swellings in the neck.

Foetal adenoma (Wolfer 1883) whether regarded as a true neoplasm arising in nests of embryonic epithelial cells or as an ordinary hyperplastic nodule in which the vesicles are unusually small and devoid of colloid (Joll 1932) presents clinically as a firm smooth single tumour within the substance of the thyroid gland. It is usually innocent.

What were believed to have been *malignant changes* were found by Wilson (1921) and by Speese and Brown (1921) in about 5 per cent of all goitres that were surgically removed but their histological criteria have been disputed and the true incidence of malignancy is probably lower. In non-toxic goitres it may be between 1 and 4 per cent (Lerman 1944) but in toxic nodular goitre it is extremely rare. Thus Means (1937) said he had not seen a single case and Crile (1936) met no instance of toxicity amongst 249 malignant cases. Malignancy should be suspected when a goitre grows rapidly, becomes unduly hard, causes dysphagia, involves the recurrent laryngeal nerve, surrounds and buries the common carotid artery, obstructs the internal jugular vein, causes pain by involving adjacent sensory nerves or when fixation can be demonstrated. Enlargement of neighbouring cervical lymph glands is particularly suggestive. Metastases are found especially in the lungs and in bone.

Riedel's disease (Riedel 1896) may be readily confused with malignant disease clinically. It is characterised by a brawny induration of part or all of the thyroid gland, sometimes involving surrounding tissues. It is a slow fibrotic process of unknown etiology affecting individuals of either sex and of any age. Pain, dyspnoea, dysphagia, huskiness of the voice and obstruction of neighbouring vessels occur and the gland is soon fixed but lymph nodes are not enlarged and thyrotoxic symptoms are unusual.

Lymphadenoid goitre (Hashimoto's disease) is seen particularly in women over the age of 45. The whole gland is involved from the start, surrounding structures are not affected and myxoedema usually develops (Joll 1932). Microscopically acinar remnants are scattered among masses of lymphoid tissue.

Acute thyroiditis may complicate a variety of infections but is rare. It may be suppurative or non suppurative according to the nature of the invading organism and to the severity of the attack. Clinically it is characterised by a painful, tender, uniform swelling of the gland accompanied by fever. Cellulitis with or without suppuration may invade surrounding tissues. Thyrotoxic symptoms may be associated but usually subside with the inflammation.

Thyroglossal cyst is essentially a mid line structure developing from remnants of the thyroglossal duct and moves upwards when the tongue is protruded. It is of cosmetic rather than medical significance.

4 *Cardiovascular signs* Vasodilatation in the skin and muscle is nearly always present and may be recognised by hot extremities, throbbing digital vessels, capillary pulsation, modified water hammer pulse and raised pulse pressure. Tachycardia is the rule and persists during sleep (Boas 1932). The action of the heart is vigorous, the cardiac impulse being forceful and displaced a little to the left, and the heart sounds slapping. A systolic murmur may be heard at apex or base, and a thrill may be felt on compressing the carotid or subclavian artery. Rarely a functional mitral diastolic murmur may be heard.

Auricular fibrillation may be initiated by overdosage of thyroxine in patients with normal hearts. It occurs in 10 per cent of all cases of thyro-

toxicosis and in 84 to 96 per cent of those with cardiac failure and may be paroxysmal or persistent. It is rare in young subjects but becomes progressively frequent with advancing years. During attacks the ventricular rate is apt to be very fast and the patient may complain of violent palpitations.

Cardiac enlargement and failure are also relatively late developments and are unusual with normal rhythm but often follow the onset of auricular fibrillation. Congestion is systemic rather than pulmonary. An appreciable proportion of such cases (over 50 per cent according to Magee and Smith 1935) are complicated by hypertension or other forms of heart disease.

X rays may show slight prominence of both the aortic knuckle and left pulmonary arc (Parkinson and Cookson 1931) and general fullness of all chambers, probably due to the high cardiac output (fig 18 05). The electrocardiogram may be within normal limits unless it shows auricular fibrillation or the voltage of P and QRS may be augmented (fig 18 06).

5 *Other and less constant features* Neurological signs are rare; they include exophthalmic ophthalmoplegia and myasthenia—sometimes resembling myasthenia gravis but not responding to prostigmine. Curious

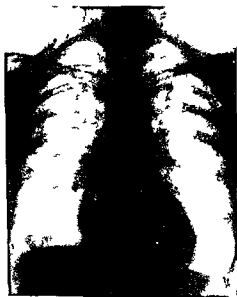


Fig 18 05—Skitag showing slight prominence of the aortic knuckle and of the left pulmonary arc in case of thyrotoxicosis.

patches of local myxœdema occasionally occur on the legs koilonychia has been described and the skin may be unduly pigmented

Decalcification of bone is not uncommon a negative calcium and nitrogen balance may be demonstrated the blood cholesterol may be rather low sugar tolerance may be reduced and impairment of hepatic function has been reported

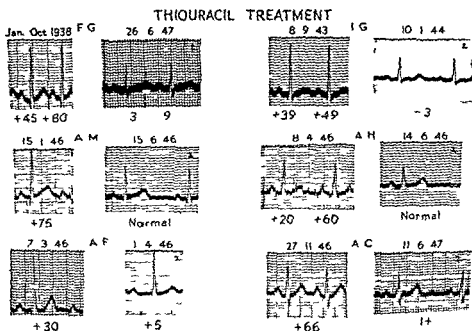


Fig. 1806—Electrocardiograms (all lead 2) showing relatively high voltage P and QRS waves in 6 cases of thyrotoxicosis. After treatment with thiouracil the voltage falls considerably. The B M R is recorded under each record

SPECIAL INVESTIGATIONS

1. *The basal metabolic rate (B M R)* introduced by Magnus Levy in 1895 has proved a useful guide to the degree of hyperthyroidism and is a measurement of the amount of oxygen consumed by the patient per minute when at complete rest i.e. fourteen hours after the last meal and after lying down undisturbed for at least half an hour. The patient breathes in and out of a closed system containing equal proportions of air and oxygen for ten minutes carbon dioxide being removed by means of soda lime the amount of gas disappearing from the system represents the total amount of oxygen consumed. This is then recorded in terms of oxygen consumption per square metre of body surface per minute and expressed as a percentage of what a normal person of the same age and sex would require. In thyrotoxicosis the B M R commonly ranges between plus 10 and plus 80 per cent. Read's formula for estimating the B M R by the pulse rate and pulse pressure is unreliable and worth no more than the knowledge that the com

bination of tachycardia and a bounding pulse suggest a raised cardiac output (Read's formula is $B M R \text{ equals } \frac{1}{3} [\text{pulse rate plus } \frac{1}{3} \text{ pulse pressure}] \text{ minus } 72$)

It should be understood that a single B M R of plus 20 per cent does not necessarily mean that the disease is milder than one with a B M R of plus 40 per cent for the course of thyrotoxicosis is variable. Serial readings may give a truer picture of the degree of activity. Another important point is that auricular fibrillation and heart failure are more often associated with low grade activity acting over a long period of time than with acute thyrotoxicosis so that the level of the B M R is no guide to the degree of cardiac disability.

The B M R is more difficult to interpret when measured for diagnostic purposes but if it is below plus 10 per cent thyrotoxicosis is improbable. High readings however may be due to faulty basal conditions or to other causes such as leukaemia and relatively high readings may be obtained in congestive heart failure of any etiology.

2 *The administration of 10 minims (0.6 ml) of Lugol's iodine* three times daily for a week or ten days may be used as a test for hyperthyroidism in two ways (1) to see whether it unmasks a goitre for a hyperplastic gland enlarges and hardens under its influence (2) to determine its effect on the sleeping pulse, body weight and B M R for these are beneficially influenced in thyrotoxic cases but not when the B M R is raised from other causes.

3 *Measurements of the cardiac output, peripheral blood flow and circulation time* provide valuable data. Outputs of 6 to 12 litres per minute are usual and are correlated more with the heart rate than with the venous filling pressure. When the heart fails the output drops usually to subnormal levels. The fore arm blood flow is invariably increased and usually remains so when the cardiac output falls as a result of failure moreover the augmented flow does not subside for several weeks after the B M R has been restored to normal by means of thyroidectomy or thiouracil therapy (Howarth 1948). Circulation times under 10 seconds are characteristic (Goldberg 1938) and may remain well within normal limits when there is systemic congestion.

The demonstration of a high cardiac output at rest places a case in the hyperkinetic group the differential diagnosis then includes severe anaemia, anoxic cor pulmonale, arterio-venous aneurysm, Paget's disease of bone, secondary carcinoma involving the liver or other serious hepatic disorders and beri beri. The majority of these can be recognised or excluded at once on clinical grounds.

4 *Urinary creatine test* Up to 200 mg of creatine may be excreted daily in the urine by normal women and children in an irregular manner but very little if any by normal men. Excessive creatinuria occurs during pregnancy and increased amounts may appear in the urine of either sex in fevers, wasting diseases and certain muscular dystrophies.

Most thyrotoxic subjects excrete an excess of creatine (Sohval, King and Reiner 1938) and its detection may be used as a diagnostic test if the above considerations are borne in mind. Thyroid responsibility may be proved by the disappearance of creatinuria within ten days of first giving iodine or thiouracil treatment (fig 18 07) (Schrire 1938). On the other

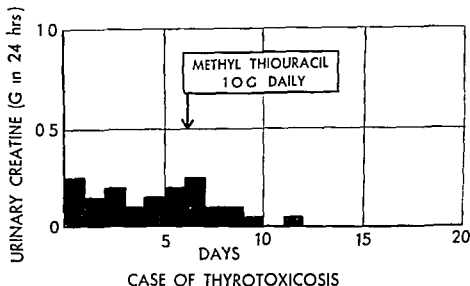


Fig 18 07—Effect of thiouracil on the excretion of creatine in the urine

hand absence of creatinuria does not exclude thyrotoxic heart disease for such cases are apt to be associated with low grade toxicity acting over a long period of time rather than with a high degree of hyperthyroidism and creatine excretion may be well within normal limits.

5 *Electrocardiography* may reveal abnormally high voltage of P and QRS (fig 18 06) as previously stated. It may also be of value in proving the nature of an irregularity of rhythm or in excluding certain other causes of a hyperkinetic circulation (e.g. pulmonary heart disease and anaemia).

6 *Radio active tracer iodine* may be used to estimate the rate at which it enters and leaves the gland (Heating *et al* 1945). In thyrotoxicosis iodine is concentrated in the gland more quickly and in greater degree than in normal controls and the excretion of iodine in the urine (after a test dose) is retarded. A Geiger counter is placed on the neck to detect the arrival and concentration of tracer iodine in the thyroid and a liquid counter is used to detect it in the urine. These tests are promising and may become of service to the clinician.

TREATMENT

The most satisfactory method of treating thyrotoxic heart disease is subtotal thyroidectomy as developed by Dunhill (1908 1929 1937). The best results are obtained when physician and surgeon work in the closest

harmony, success depending as much upon the skill and judgment of the physician as upon the experience and dexterity of the surgeon (Fraser and Dunhill 1934) adequate premedication being all important

The patient should be put to bed and fed on a liberal and nourishing diet. The addition of 5 to 10 mg. of aneurin daily may be helpful on the grounds that an abundant supply of this vitamin is needed for the increased carbohydrate metabolism. Fatigue and weakness may respond to 50 mg. of pyridoxine daily (Soskin and Levine 1944). Phenobarbitone $\frac{1}{2}$ a grain (32 mg.) t.d.s. or potassium bromide 10 grains (0.64 G.) t.d.s. may also be prescribed with benefit and a nocturnal sedative is usually necessary.

During this preliminary stage of treatment which usually induces some remission of symptoms the degree of thyrotoxicosis may be assessed clinically and by means of the special tests detailed above. Iodine may then be given by mouth in doses of 10 minims (0.6 ml.) of Lugol's solution three times daily preferably in milk. Within ten days there is usually marked improvement the pulse rate falls the B.M.R. is lowered and the patient feels better (Waller 1914 Plummer 1923). The moment for operation is usually ten to fourteen days after beginning iodine. If however the patient is not ready at that time there should be no hesitation in postponing it but the dose of iodine should then be reduced to 5 minims (0.3 ml.) three times a day (Fraser and Dunhill 1934).

The introduction of thiouracil by Astwood (1943) following the discovery by the Mackenzies (1941) that the administration of sulphaguanidine to rats caused thyroid hyperplasia and reduction of colloid has proved an important therapeutic advance. Thyroid hyperplasia was attributed to increased production of thyrotropic hormone by the anterior pituitary in an endeavour to compensate for deficiency of thyroid hormone brought about by sulphaguanidine. Astwood found that many substances had a similar effect including all the sulphonamides *p*-aminobenzoic acid thiourea and its compounds and that of these thiouracil offered the best prospects being potent and relatively non-toxic. It is held that thiouracil and the other substances mentioned act by interfering with the union of iodine and tyrosine and so prevent the formation of di-iodotyrosine a known precursor of thyroxine (Riker and Wescoe 1945). The histological appearance of the thyroid gland under their influence resembles the hyperplastic gland of iodine deficiency.

Extensive trials have established the dosage toxic effects and early results of thiouracil treatment on firm ground (Astwood 1944 Williams 1944). Reduction of the B.M.R. and amelioration of all symptoms except exophthalmos and those due to pressure from the goitre are obtained whether the hyperthyroidism is due to primary Graves' disease or to toxic nodular goitre. The usual dose is 0.2 G. of thiouracil twice daily for about three weeks or until the available evidence suggests that the production of thyroid hormone has been reduced to normal levels. During this stage the white blood cells may be counted weekly and the patient is

best confined to bed in hospital. If medical treatment is continued the dose is then reduced to 0.2 G daily for a month or so, and the patient is allowed to resume her normal activities. Next the maintenance dose is discovered by trial and error: it varies considerably from case to case but is of the order of 0.05 to 0.2 G daily. According to Himsworth (1948) equally good results are obtained when the initial dose is only 200 mg daily, and the maintenance dose 50 to 100 mg. Himsworth claims that the results of thiouracil therapy are as good as those of subtotal thyroidectomy and that medical treatment should therefore be preferred because the mortality rate is lower. Auricular fibrillation may revert to normal rhythm spontaneously with thiouracil treatment or normal rhythm may be restored by means of quinidine. Treatment should be continued for at least twelve months if medical cure is desired but even then 33 to 50 per cent of cases relapse when the drug is withdrawn (Himsworth 1948; Williams 1946).

Signs and symptoms of myxœdema may develop if too much thiouracil is given but soon disappear when the dose is reduced. Toxic reactions occur in 13 per cent of cases (Van Winkle 1946) usually during the first three weeks and include nausea and vomiting, sulphonamide-like rashes, fever, agranulocytosis, purpura and adenopathy. The most serious of these is agranulocytosis (1 to 2 per cent) which should be treated promptly with penicillin, pentnucleotide, liver extract and perhaps blood transfusion, thiouracil being abandoned. An uncontrollable hæmorrhagic state ending in renal failure was the cause of death in one case known to the author. The mortality rate has been about 0.5 per cent (Moore 1946) but may be less now that physicians are more experienced in using the drug. Aminothiazol 0.2 to 0.8 G daily (Perrault 1946) has been used widely in France but has little advantage over thiouracil. Methyl thiouracil in similar doses is less toxic and propyl thiouracil least so; both are as potent as thiouracil (Astwood and VanderLaan 1945).

In view of the high relapse rate on discontinuing thiouracil treatment, and because the operative risk and difficulties may then be greater owing to the increased vascularity of the gland, most workers favour the drug mainly as a preliminary to partial thyroidectomy. Moreover thiouracil has not replaced iodine in this respect for the subsequent administration of Lugol's solution is found to diminish the vascularity of the gland and so to facilitate the operation (Means 1946). Again in most cases of primary Graves' disease pre-operative iodine is so satisfactory that thiouracil is hardly justified. But in severely toxic cases and in most instances of toxic nodular goitre with or without cardiac embarrassment thiouracil is superior to iodine. They may be given together with advantage in doses of 0.1 G – 0.2 G of methyl or propyl thiouracil and 5 minims (0.3 ml) of Lugol's iodine three times daily. Symptoms should be controlled within two weeks. Subtotal thyroidectomy may then be carried out when convenient: there is no urgency as there is when iodine is used alone because patients do not relapse while taking sufficient thiouracil.

Cardiac complications do not contraindicate partial thyroidectomy (Dunhill 1937) More careful preparation however is needed auricular fibrillation must be controlled and heart failure relieved before it is safe to operate but normal rhythm should not be deliberately restored at this stage

The commonest post operative complication has been paroxysmal auricular fibrillation with rapid ventricular rate but this may be less frequent if the patient is prepared with thiouracil It should not occasion undue alarm for the rhythm usually reverts spontaneously to normal within 48 hours If auricular fibrillation persists however whether previously well established or of recent onset every effort should be made to restore normal rhythm by means of quinidine before the patient leaves hospital (see page 149) The risk of embolism is slight perhaps because the hyperkinetic circulation lessens the chance of venous thrombosis

More recently attempts have been made to treat toxic goitre with a combination of thiouracil and thyroxin It has been pointed out that exophthalmos and the size and vascularity of the goitre may be increased by thiouracil and have been attributed to the liberation of increased quantities of thyrotropic substance owing to deficiency of thyroid hormone Although hyperthyroidism obviously cannot result from this thyrotropic activity proper cure of the disease may well be frustrated by its presence Moreover the vascularity of the gland may become so great that it may function as an arteriovenous aneurysm and so maintain the hyperkinetic circulation thiouracil was meant to relieve A continuous thrill and loud machinery murmur may then be appreciated over the gland In one such case investigated by Wyndham and the author admittedly the result of overdosage the resting cardiac output was twelve litres per minute But if a maintenance dose of thyroid (1 to 3 grains or 60 to 180 mg daily) is given in conjunction with thiouracil pituitary hyper activity may be prevented or may subside if already present (Williams and Bissell 1943) It is possible that medical cure may be more readily achieved along these lines

Three other methods of treatment may be mentioned 1 *X ray therapy* is curative in about a third results in some improvement in a third and is without benefit in the remainder (Means and Holmes 1923) It may be useful in thiouracil sensitive subjects who refuse operation 2 *Pituitary irradiation* has been tried with limited success (Thompson and Thompson 1944) but not very widely Interference with other pituitary functions is the obvious disadvantage even if greater antithyrotropic success were achieved 3 Preliminary reports on treatment with *radio active iodine* have been favourable (Hertz and Roberts 1942-6) As the substance is largely concentrated in the thyroid gland the irradiation effect is considerable but the difficulty in gauging the correct dose and the danger of exciting malignant changes in the gland are possible drawbacks

Thyrototoxic crises Owing to the impossibility of neutralising thyroid hormone that has already been manufactured both iodine and thiouracil do

not benefit the patient for several days (graphs illustrating the effect of partial thyroidectomy, iodine and thiouracil on the basal metabolic rate are remarkably similar). The treatment of thyrotoxic crises by massive doses of iodine (by mouth or intravenously) as advocated by Boland and Kepler (1938) for example is therefore questionable. Absolute rest, heavy sedation and replacement of salt and water lost in sweating and vomiting, are probably more important. Aneurin, 100 mg intravenously may also help.

If toxic goitre is recognised and treated promptly however crises should not occur.

Thyrotoxicosis and tonsillitis Cases are encountered in which an attack or repeated attacks of tonsillitis are associated with thyrotoxicosis. The problem then arises whether to perform partial thyroidectomy or tonsillectomy first. Before the introduction of thiouracil most authorities agreed that it was safer to remove the thyroid gland before the tonsils for tonsillectomy in thyrotoxic patients sometimes precipitated a crisis. Thiouracil has simplified the problem however and allows tonsillectomy to be undertaken first without risk.

Thyrotoxicosis and rheumatic heart disease Thyrotoxicosis may be associated with acute rheumatic carditis or with established rheumatic valve lesions. Both Parry's and Basedow's first cases were so related. The association if more than a coincidence is indirect and may depend upon their joint relationship to streptococcal tonsillitis. Treatment aims at partial thyroidectomy as soon as the rheumatic state allows it. Rheumatic heart disease with fixed valve lesions may result in enormous enlargement of the heart owing to the excessive work induced by thyrotoxicosis (fig. 1808) and the sooner the latter is treated the better. Total thyroid ablation however is not indicated.

Thyrotoxicosis and hypertension There is a group of cases sometimes designated thyrotoxic hypertension in which thyrotoxicosis is associated with high blood pressure both systolic and diastolic levels being raised. There is little evidence of any direct relationship between the two diseases and the blood pressure does not fall following thyroidectomy (Bisgard 1939).



Fig. 1808—X-ray showing gross cardiac enlargement in a case of thyrotoxicosis plus mitral stenosis.

Thyrotoxicosis and angina pectoris Ischæmic heart pain occurs when the blood supply to the myocardium is insufficient to meet the demand. By increasing the demand thyrotoxicosis may induce angina in a patient with a relatively minor degree of coronary atherosclerosis behaving in this respect like anæmia. Thyroid hormone also sensitises the organism to adrenalin. When ischæmic and thyrotoxic heart disease are associated subtotal thyroidectomy need not be withheld on the grounds of undue risk for the operation may be followed by many years of normal life before angina again makes its appearance.

Thyrotoxicosis and pregnancy Thyrotoxicosis developing during pregnancy may be due to primary exophthalmic or nodular goitre. With the aid of thiouracil in combination with small doses of iodine or thyroid patients should be taken safely to term. If the condition does not then subside subtotal thyroidectomy may be carried out. The danger of goitre developing in the fœtus is minimised by the iodine (or thyroid) but it is well to keep the dose of thiouracil as small as possible (not more than 0.2 G daily).

PROGNOSIS

There are few forms of heart disease that respond better to adequate treatment than thyrotoxic heart disease. Cases with gross congestive failure and well established auricular fibrillation may be cured and the largest hearts may resume their normal size (fig. 18.09). On the other hand heart failure and death are inevitable if the disease remains unchecked. In the



Fig. 18.09 (a)—Thyrotoxic heart failure



(b)—After subtotal thyroidectomy

hands of the best surgeons the mortality rate of subtotal thyroidectomy in cases of toxic nodular goitre has been 1.6 per cent (Cole 1944) to 2.6 per cent (Dunhill 1937) but it may be less with thiouracil preparation. No reliable figures are available upon which to assess the total relapse rate. Post-operative tetany and paralysis of the vocal cord each occurs in approximately 1 per cent (Means 1946).

According to Himsworth (1948) there is a 3:1 chance in favour of a permanent remission with thiouracil treatment.

THE HEART IN MYXŒDEMA

Artificial myxœdema produced by total ablation of the thyroid gland or by thiouracil benefits the heart by lessening the circulatory demands and so relieves angina pectoris and congestive heart failure. Yet well developed myxœdema from natural causes gives rise to cardiac enlargement, pericardial effusion and ultimately to congestive heart failure; moreover angina pectoris may be associated. Enlargement cannot be due to overwork; it must depend upon some intrinsic change in the heart muscle. Histological examination however is usually disappointing. The fault is probably biochemical and is unlikely to be properly understood until studies in tissue chemistry are more advanced.

The diagnosis of myxœdema is suggested by the placid sleepy character (unless there is manic psychosis), poor memory, sensitivity to cold (Raynaud's phenomenon is common), dry coarse skin, thickened lips and tongue, low thick voice, baggy eyes, scanty dry hair, podgy hands, supraclavicular pads of fat and general pallor. It is confirmed by an impalpable thyroid gland, by a B.M.R. of minus 30 to 40 per cent, by prolongation of the arm to tongue circulation time to 19 to 25 seconds, by a high blood cholesterol of 300 to 400 mg. per cent, by relative insensitivity to atropine and adrenaline, by a characteristic form of anæmia and by a pathognomonic electrocardiogram.

The type of anæmia which responds to thyroxine alone is normocytic and orthochromic and may be regarded as a compensatory adjustment to diminished oxygen requirement (Bomford 1938). The electrocardiogram shows sinus bradycardia, low voltage auricular and ventricular complexes and flat or inverted T waves in all leads (fig. 18.10). The cause of these changes is not yet understood; they do not depend upon the presence of pericardial effusion nor upon the state of the subcutaneous tissues. The response to thyroxine is quick and complete and accompanies beneficial changes in the B.M.R. The electrocardiogram in cretinism behaves similarly (fig. 18.11).

Whilst a well developed case of myxœdema is difficult to overlook (fig. 18.12) cases of short duration, especially in younger women (the sex incidence is 8:1 in favour of women) may easily escape notice. The diagnosis should be considered in any case of congestive heart failure or of peri-

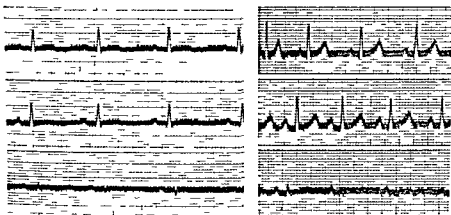


Fig 18 10 (a)—Electrocardiogram showing sinus bradycardia low voltage auricular and ventricular complexes and flat T waves in all leads in a case of myxœdema (b) Normal electrocardiogram after treatment

cardial effusion of unknown etiology Congestion when it occurs is systemic and is associated with a low cardiac output Pericardial effusion is due to simple transudation Cardiac enlargement is general (fig 18 13) and pulsation of all chambers is poor Angina pectoris is said to occur in 1 to 2 per cent of cases (Smyth 1938) but is probably more frequent Coronary atherosclerosis may result from the high blood cholesterol

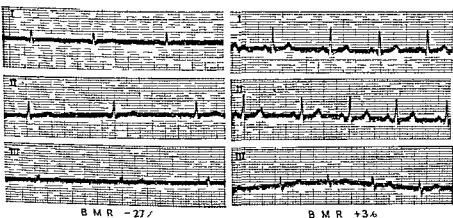


Fig 18 11—Electrocardiogram before and after treatment in a case of cretinism

Myocardial infarction without coronary thrombosis has been described in such cases when treated too vigorously with thyroxine The blood pressure is little influenced by myxœdema and is as often elevated as low When congestive failure is present measurements of the BMR give unduly high readings more reliance should then be placed on other tests especially on the electrocardiogram



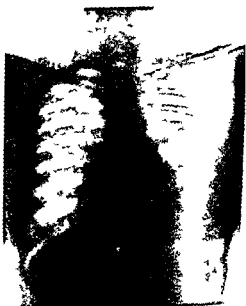
Fig. 1813 (a) — Before treatment



(b) After seven weeks treatment



Fig. 1813 (a) — Radiogram showing enlargement of the carcinoma before treatment



(b) After treatment

Treatment If there is no evidence of coronary disease thyroxine may be given intravenously in a single dose of 10 mg or thyroid may be given by mouth in dose of 3 grains (0.2 G) daily. The response is delayed but dramatic. Within five to ten days the B.M.R. rises, the blood cholesterol falls, I wave changes are corrected and clinical improvement is obvious. Signs of failure or of pericardial effusion soon disappear and the heart resumes its normal size (Ierman, Clark and Means, 1933).

Initial treatment is easier than maintenance. With the aid of the B.M.R. it is not difficult to regulate dosage for a patient at rest in bed, but when she leaves hospital and varies her activities it is not so easy and supervision is required for life. The average maintenance dose of thyroid extract is 2 to 3 grains (0.13 to 0.2 G) daily by mouth.

If there is any suspicion of associated coronary disease initial treatment should be cautious and the oral route advised. Not more than 1 grain (64 mg) of thyroid extract should be given daily and in cases with angina pectoris not more than 1/2 a grain (32 mg). The dose may be increased slowly week by week if well tolerated or reduced and maintained at a minimum if not tolerated.

The chief complication arising during treatment is the development of angina pectoris; should this occur the dose of thyroid may have to be less than ideal but enough to keep the blood cholesterol below 300 mg per cent.

REFERENCES

- Astwood E. B. (1943) Treatment of hyperthyroidism with thiourea and thiouracil. *J. Amer. med. Ass.* 122: 78. — (1944-5) Chemotherapy of hyperthyroidism. The Harvey Lectures series 40: 195. — VanderLaan W. P. (1945) Thiouracil: derivation of greater activity for treatment of hyperthyroidism. *J. clin. Endocrinol.* 5: 424.
- von Basedow C. A. (1849) Exophthalmos durch Hypertrophie des Zellgewebes in der Augenhöhle. *Monatsschrift für die gesammte Heilkunde* Berlin 28th March.
- Baumann E. (1895) Ueber das normale Vorkommen von Jod im Tierkörper. *Z. f. physiol. Chem.* 21: 319.
- Bisgard J. D. (1939) Relation of hyperthyroidism to hypertension. *Arch. intern. Med.* 63: 497.
- Boas E. P. (1932) Heart rate during sleep in Graves' disease and in neurogenic sinus tachycardia. *Amer. Heart J.* 8: 4.
- Boland E. W. and Kepner F. J. (1938) Crisis of exophthalmic goitre. Report of case. *Proc. Mayo Clin.* 13: 817.
- Bornford R. R. (1938) Anæmia in myxœdema and role of thyroid gland in erythropoiesis. *Quart. J. Med.* 7: 495.
- Brain W. R. (1939) Exophthalmos in Graves' disease despite sympathetic paralysis. *Lancet* ii: 1217. — Turnbull N. M. (1938) Exophthalmic ophthalmoplegia with pathological report on ocular muscles and thyroid glands. *Quart. J. Med.* 7: 93.
- Bruun E. (1945) Exophthalmic goitre developing after treatment with thyroid preparation. *Acta med. Scand.* 122: 13.
- Cole W. H. (1944) Factors influencing operability and mortality rate in goitre. *Surg.* 16: 688.
- Crisle G. Jr. (1936) Hyperthyroidism associated with malignant tumours of the thyroid gland. *Surg. Gynec. and Obstet.* 62: 995.
- Dunhill T. (1908) The surgical treatment of exophthalmic goitre. *Intercol. med. J. Australia* 13: 793. — (1929) Toxic goitre. *Brit. J. Surg.* 17: 4-4.
- (1937) Surgery of the thyroid gland. The Lettsomian lectures. London.

- Flajani G (1802) Sopra un tumor freddo nell' anterior parte del collo detto broncocele *J* 270 Rome
- Fraser F R and Dunhill T P (1934) Lectures on toxic goitre London
- Friedberg C K and Sohval A R (1937) Occurrence and pathogenesis of cardiac hypertrophy in Graves' disease *Amer Heart J* 13 599
- Goldberg S J (1938) Circulation time as diagnostic aid in hyperthyroidism *Ann intern Med* 11 1818
- Graves R J (1835) Newly observed affection of the thyroid gland in females *London med and surg J* 7 516
- Harrington C R (1933) The thyroid gland London ——— Barger G (1927) Chemistry of thyroxine II Constitution and synthesis of thyroxine *Biochem J* 21 169
- Hertz S and Roberts A (1942) Application of radioactive iodine in therapy of Graves' disease *J clin Invest* 21 624 ——— (1946) The use of radioactive iodine therapy in hyperthyroidism *J Amer med Ass* 131 81 ———
- Evans R D (1938) Radio active iodine as indicator in the study of thyroid physiology *Proc Soc exper Biol and Med* 38 510
- Humsworth H P (1948) Thiouracil and its derivatives in the routine treatment of thyrotoxicosis *Brit med J* 11 61
- Howarth S (1948) Personal communication
- Joll C A (1932) Diseases of the thyroid gland London
- Keating F R Rawson R W Peacock W and Evans R D (1945) Collection and loss of radio active iodine compared with anatomic changes induced in thyroid of chick by injection of thyrotropic hormone *Endocrinol* 36 137
- Kendall E C (1915) The isolation in crystalline form of the compound containing iodine which occurs in the thyroid its chemical nature and physiologic activity *J Amer med Ass* 64 2042
- Kepler E J and Barnes A R (1932) Congestive heart failure and hyperthyroidism in hyperthyroidism Clinical and pathological study of 178 fatal cases *Amer Heart J* 8 102
- Lerman J (1944) The endocrine activity of thyroid tumours and the influence of the thyroid hormone on tumours in general *Surg* 16 266 ——— Clark R J and Means J H (1933) Heart in myxoedema electrocardiograms and roentgen ray measurements before and after therapy *Ann intern Med* 6 1251
- Magee H R and Smith H L (1935) Auricular fibrillation in hyperthyroidism influence of age *Amer J med Sc* 189 683
- Magnus Levy A (1895) Ueber den respiratorischen Gaswechsel unter dem Einfluss der Thyreoidea sowie unter verschiedenen pathologischen Zuständen *Berl klin Hoch* 32 650
- Marie P (1883) Sur la nature et sur quelques uns des symptomes de la maladie de Basedow *Arch de Neurol* 6 79
- Marine D (1927) Iodine in the treatment of diseases of the thyroid gland *Medicine* 6 127 ——— (1930) The essential thyroid changes in goitre *Amer J Path* 6 607 ——— Lenhart C H (1910) Observations and experiments on the so called thyroid carcinoma of brook trout (*salvelinus fontinalis*) and its relation to ordinary goitre *J exper Med* 12 311 ——— Rosen S H (1933) Exophthalmos in thyroidectomized guinea pigs by thyrotropic substance of anterior pituitary and the mechanism involved *Proc Soc exper Biol and Med* 30 901
- McCarrison R (1927) Experiment in goitre prevention *Brit med J* 1 94
- Mackenzie J B Mackenzie C G and McCollum F V (1941) *Science* 94 518
- Means J H (1937) The thyroid and its diseases Philadelphia ——— (1946) Evaluation of the several methods for treating Graves' disease available to day *Ann intern Med* 25 403 ——— Holmes G W (1923) Further observations on the Roentgen ray treatment of toxic goitres *Arch intern Med* 31 303 ———
- Richardson E P (1938) The diagnosis and treatment of diseases of the thyroid New York
- Moebius P J (1886) Ueber Insufficienz der Konvergenz bei morbus Basedowii *Centralbl f nerventk u Psychiat* 9 356

- Moore F D (1946) Toxic manifestations of thiouracil therapy a co operative study *J Amer med Ass* 130 315
- Oswald A (1899) Die Eiweisskörper der Schilddrüse *Z f physiol chem* 27 14
- Parkinson J and Cookson H (1931) Size and shape of heart in goitre *Quart J Med* 24 499
- Parry C H (1825) Collected works 1 478 London (Extracted by Major R H in Classic descriptions of disease Springfield Illinois 1932)
- Perrault M (1946) Aminothiazol (2921 R P) in the treatment of Graves disease *Paris Med* 36 401
- Plummer H S (1923) Results of administering iodine to patients having exophthalmic goitre *J Amer med Ass* 80 1925
- Pochin E E (1937-8) Unilateral retraction of upper lid in Graves disease *Clin Sc* 3 197 — (1939) Ocular effects of sympathetic stimulation in man *Ibid* 4 79 — (1939) Mechanism of lid retraction in Graves disease *Ibid* 4 91
- Rake G and McEachern D (1932) Study of heart in hyperthyroidism *Amer Heart J* 8 19
- Riedel (1896) Die chronische zur Bildung eisenharter Tumoren führende Entzündung der Schilddrüse *Verhandl d deut Ges f Chir* 25 101
- Riker W F and Wescoe W C (1945) The pharmacology and therapeutic application of anti thyroid compounds *Amer J med Sc* 110 665
- Schrire I (1948) The effect of 2 thiouracil on the creatinuria of thyrotoxicosis and its use in the diagnosis of thyrotoxicosis *Clin Sc* 7 49
- Shaw R C (1909) Cervical sympathetic and its relation to thyroid gland in exophthalmic goitre *Brit med J* 1 495
- Smyth C J (1938) Angina pectoris and myocardial infarction as complications of myxœdema *Amer Heart J* 15 652
- Sohval A R King F H and Rainer M (1938) The creatine tolerance test in the diagnosis of Graves disease and allied conditions *Amer J med Sc* 195 608
- Soskun S and Levine R (1944) Recent advances in physiology of the thyroid and their clinical application *Arch intern Med* 74 375
- Speese J and Brown H P Jr (1921) Malignant degeneration in benign tumours of the thyroid gland *Ann Surg* 74 684
- Thompson W O and Thompson P K (1944) Treatment of toxic goitre by irradiation of the pituitary *J clin Invest* 23 951
- von Graefe A (1864) Concerning Basedow's disease *Deutsch Klinik* 16 158
- van Winkle W (1946) Clinical toxicity of thiouracil survey of 5 745 cases *J Amer med Ass* 130 343
- Waller H E (1914) On the value of iodine taken internally in Graves disease *Prescriber* 8 153
- Williams R H (1944) Antithyroid drugs with particular reference to thiouracil *Arch intern Med* 74 479 — (1946) Thiouracil treatment of thyrotoxicosis I Results of prolonged treatment *J clin Endocrinol* 6 1
- Williams R and Bissell J (1943) Treatment of hyperthyroidism with thiouracil *New Engl J Med* 229 97
- Wilson L B (1911) Malignant tumours of the thyroid *Ann Surg* 74 129
- Wolffler A (1883) Über die Entwicklung und den Bau des Kropfes *Arch f klin Chir* 29 1 754
- Zondek H and T'cho A (1945) Observations on so called thyrotropic exophthalmos *Brit med J* 1 836

CHAPTER XIX

HYPERKINETIC CIRCULATORY STATES

(ANÆMIA PREGNANCY ARTERIO VENOUS ANEURYSM
BERI BERI PAGET'S DISEASE OF BONE HEPATIC
FAILURE)

IN addition to the diseases enumerated above hyperkinetic circulatory states (Harrison 1935) include thyrotoxicosis anoxic pulmonary heart disease fever and exercise. The first two have been considered fully elsewhere and the last two have a purely physiological basis.

All these conditions are characterised by a raised cardiac output maintained by means of tachycardia a raised venous filling pressure or both moreover the heart may beat more strongly. Conspicuous evidence of vaso-dilatation in skin and muscle is found in all of them the skin is warm and flushed the pulse is collapsing the digital vessels throb and there may be capillary pulsation in fact the peripheral circulation resembles that seen in aortic incompetence. The fore arm and calf blood flows are also increased. Whilst young and healthy hearts may cope with the situation without distress older or unhealthy hearts may fail to meet the requirements.

It may be difficult clinically to recognise congestive failure in these cases for the usual signs may have other interpretations. Thus a raised venous pressure may be part of the physiological mechanism maintaining a high cardiac output (McMichael 1947) enlargement of the liver may be due to secondary carcinoma or to hepatitis and œdema is commonplace in severe anæmia and beri beri for other reasons. Indeed it is by no means easy to be sure what is meant by failure in this group for example McMichael uses the term high output failure to describe a state in which a raised venous pressure and œdema are associated with a high cardiac output whether or not the latter is capable of being raised further. Yet failure ordinarily denotes an overloaded heart or ventricle one incapable of raising its output further. But this question has already been discussed (page 155).

THE HEART IN ANÆMIA

Physiology. Severe chronic or post hæmorrhagic anæmia may affect the heart in three ways (i) it may cause a hyperkinetic circulatory state as already described (ii) it may cause or precipitate angina pectoris or acute coronary insufficiency (iii) it may result in nutritional degenerative changes in the cardiac muscle which may reduce its reserve.

With an oxygen consumption of 240 ml per minute an anæmic subject with a hæmoglobin of 20 per cent could not have a cardiac output less than 6 litres per minute if all the available oxygen were utilised (0 per cent Hb = 3 G Hb per cent = 3×1.34 ml oxygen per cent = 4 ml oxygen per cent or 40 ml per litre. Thus cardiac output = $\frac{240}{40}$ = 6 litres per minute).

If half the available oxygen were utilised the cardiac output would be 12 litres per minute.

In anæmic subjects investigations have shown that the resting cardiac output may reach 13 litres per minute and utilisation of available oxygen may be increased from the normal 33 per cent to as much as 90 per cent (Liljestrand and Stenstrom 1925-6, Nielson 1934, Sharpey-Schuler 1944). These changes do not occur at rest with hæmoglobin values above 50 per cent but become increasingly apparent at lower levels (Bouchut and Froment 1934). The high cardiac output is maintained both by tachycardia and a raised venous pressure. The latter must be due to widespread capillary or peripheral venoconstriction for the blood volume is reduced (McMichael *et al.* 1943) and the small arteries and arterioles are dilated (McMichael 1947).

Clinical features. The chief symptoms of severe anæmia are breathlessness, fatigue and palpitations. Angina pectoris occurs in about 30 per cent (Coombs 1926, Pickering and Wayne 1934) occasionally even when there is no underlying coronary disease. Thus the author has treated a boy of 17 with pernicious anæmia and angina pectoris and also a young man of 21 who presented himself with classical ischæmic heart pain due to iron deficiency anæmia resulting from bleeding hæmorrhoids. Oedema may be due to congestive heart failure but is more often nutritional. It is especially prone to develop during the first three weeks of blood regeneration in response to treatment of the anæmia.

Paroxysmal cardiac dyspnoea or acute pulmonary oedema is rare as a spontaneous event but may arise during blood transfusion or saline infusion. These procedures should not be lightly undertaken in cases of severe chronic or post hæmorrhagic anæmia. Precautionary measures include the use of concentrated red cells instead of whole blood and venous pressure lowering agents such as cuffs applied to the thighs. Transfusion should be temporarily abandoned if the venous pressure is seen to rise appreciably.

Physical signs. A hyperkinetic circulation and peripheral vasodilatation may be recognised by the features detailed previously.

A functional systolic murmur (so called hæmic murmur) at apex or base is common and is probably due to mitral incompetence resulting from left ventricular dilatation or to the increased velocity of blood flow. Functional mitral or aortic diastolic murmurs may also be heard occasionally, earlier observations such as those by Von Noorden (1891), Sahli (1893) and Kraus (1905) having been amply and repeatedly confirmed (Goldstein and Boas

1927) Mitral presystolic or diastolic murmurs are probably due directly or indirectly to the increased velocity of blood flow the mechanism being almost certainly the same as that responsible for mitral diastolic murmurs in patent ductus arteriosus ventricular septal defect and thyrotoxicosis. Basal diastolic murmurs are probably due to dilatation of the aortic or pulmonary rings.

The electrocardiogram Despite several publications emphasising the normality of the electrocardiogram in anaemia (e.g. Smith 1933 Pickering and Wayne 1934) there can be no doubt that significant changes occur in at

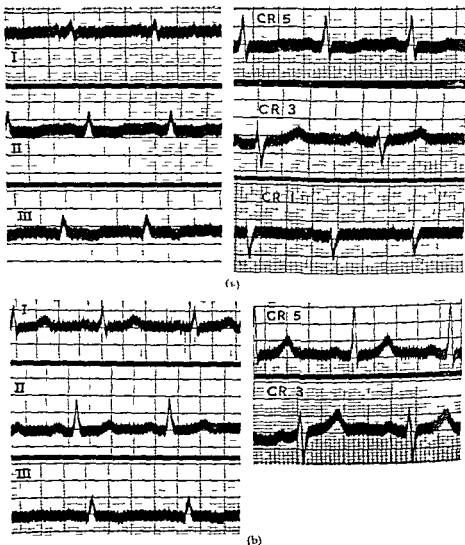


Fig 19.01—Electrocardiogram showing low voltage and flat or inverted T wave in all leads in a case of pernicious anemia

(a) Before treatment

(b) After correction

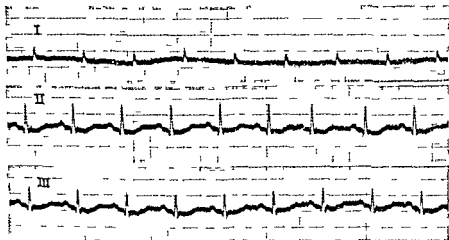


Fig 19 02—Electrocardiogram showing depression of the ST segment due to acute coronary insufficiency resulting from post hæmorrhagic anæmia



(a) Before treatment



(b) After treatment of the anæmia

Fig 19 03—Ski gram showing general cardiac enlargement in a case of severe pernicious anæmia

least a third of cases with hæmoglobin values under 40 per cent (Block 1937) In a consecutive series of twenty such cases analysed by the author eight showed low voltage depressed S T segments or flat or inverted T waves in left ventricular surface leads or their equivalents As the anæmia improved under treatment these faults were corrected (fig 19 01) Several instances of bundle branch block have also been observed but these have always persisted when the anæmia was cured Depression of the S T segment is common following gross hæmorrhage and is believed to represent temporary coronary insufficiency (fig 19 02)

Fluoroscopy X-rays often reveal slight enlargement of all chambers of the heart and prominence of both the aorta and pulmonary artery in cases with hæmoglobin levels below 40 per cent (fig 19 03)

Necropsy studies have revealed slight increase of heart weight (350 to 450 G) in the majority of cases of severe anæmia and considerable increase occasionally (Cahot and Richardson 1919) Experimental anæmia in rats has resulted in slight cardiac hypertrophy at hæmoglobin levels of 10 G per cent and considerable hypertrophy (weight at least twice normal) at levels of 2 to 3 G per cent (Forman and Daniels 1930-1) According to Grunberg (1930) hypertrophy is invariable in man when the hæmoglobin is 15 per cent or less and does not occur at all when the hæmoglobin is 66 per cent or more

These findings harmonise with the behaviour of the cardiac output in relation to hæmoglobin levels and there can be little doubt that enlargement depends on increased work

Clinical diagnosis Knowledge of cardiovascular behaviour is of little value in making a diagnosis of anæmia and is of no value at all in determining the nature of the anæmia It is helpful, however, in differential diagnosis especially between anæmia the anxiety states and bacterial endocarditis Thus an anxiety state may present with the same group of symptoms including pallor and there may be cardiac over action and functional systolic murmurs The pallor however is due to peripheral vasoconstriction and does not affect the conjunctivæ or the mucous membranes and it is less obvious in the palms of the hands the nail beds too are more likely to be cyanosed than pale In anæmia pallor is often waxy chalky or lemon tinted according to its severity and type The cardiovascular dynamics are quite different Over action of the heart and tachycardia in the anxiety states are associated with little or no rise in cardiac output there is peripheral vasoconstriction rather than vasodilatation and the diastolic blood pressure tends to be raised in casual readings the stroke volume tends to be reduced and the pulse may be small the circulation and venous pressure are normal

A type of case that may cause confusion is one that presents with pallor low grade fever petechiæ splenomegaly over action of the heart and a loud systolic murmur at apex or base Bacterial endocarditis may be suspected especially when there is a diastolic basal murmur as well and the

pulse is collapsing yet all these features may be due to anæmia alone

Treatment All cardiovascular changes due to anæmia are reversible if the anæmia is treated successfully. Cardiac remedies are rarely required apart from urgent measures in the event of acute pulmonary œdema (page 190). The danger of ill judged or too rapid intravenous infusion has already been mentioned.

THE HEART IN PREGNANCY

Physiology Changes in the cardiovascular system during normal pregnancy probably depend upon increased metabolism raised intra abdominal pressure increase in blood volume and upon the presence of a uterine

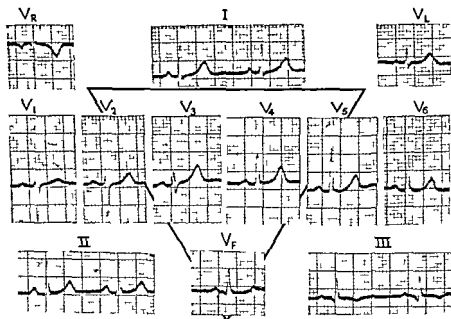


FIG. 19.04—Electrocardiogram showing characteristic appearances associated with pregnancy

arterio venous shunt. The oxygen consumption increases by about 20 per cent, the cardiac output by about 50 per cent, the arterio venous oxygen difference is decreased and the pulse rate, pulse pressure and venous pressure increased (Burwell *et al.* 1938). Faintness in the supine position may be due to pressure of the uterus on the inferior vena cava so that blood is dammed back in the legs with a resulting fall in right auricular pressure. Hot tingling extremities may be a reflection of increased peripheral blood flow.

Less important findings include functional systolic murmurs, ectopic beats, accentuation of the third heart sound, displacement of the apex beat

to the left and upwards slight increase in the size of the heart occasional minimal œdema and a characteristic electrocardiogram showing a prominent S_1 and Q_3 inversion of T_3 (fig 19 04) and a horizontal frontal plane vector

The changes described may begin as early as the second month reach their maximum at about the end of the sixth month and are maintained to term

The normal heart copes with the increased burden of pregnancy without difficulty but diseased hearts may be in grave distress The chief conditions that may give rise to anxiety are those that may affect women of child bearing age namely congenital rheumatic thyrotoxic and hypertensive heart disease and bacterial endocarditis

CONGENITAL HEART DISEASE

This is not hereditary so there is no danger of its transmission to the fœtus

Patent ductus arteriosus and patent interventricular septum unless gross are no bar to pregnancy Coarctation of the aorta usually causes little trouble and is chiefly of interest because the associated hypertension may be mistaken for toxæmia of pregnancy occasionally however the aorta has ruptured Atrial septal defect requires watching Relatively mild cases may be taken to term safely but if right sided enlargement is advanced pregnancy should be prevented or should be terminated in the first three months if possible The rise in right auricular pressure may cause reversal of the shunt and sudden severe cyanosis or the increasing stress may precipitate right ventricular failure which may also cause reversal of the shunt Pulmonary stenosis is usually a direct bar to pregnancy unless mild and of simple valvular type

RHEUMATIC HEART DISEASE

There are some who maintain that any woman who has rheumatic heart disease should be advised against having any children They argue that pregnancy affects her adversely that the strain of bringing up children shortens her life and that her premature death leaves her family stranded at a critical time Others feel that to forfeit so much human happiness on these grounds is both undesirable and unnecessary Is life so precious to prolong if so much of its meaning is taken away? Moreover available modern statistics barely support the first argument Thus in four combined series collected by Jensen (1938) the average age of death in spinsters or nulliparous women with mitral stenosis was 36 6 in married women with families it was 40 3 Again Bunim and Rubricius (1948) could find no significant difference in the life histories of 169 rheumatic mothers and 215 rheumatic childless women Of course the childless women may have been advised against pregnancy owing to the severity of their condition so that the two groups may not be strictly comparable there is insufficient

evidence on this point. It is certain however that many women with mitral stenosis unaware that there is anything wrong with them have large families and lead normal lives until the lesion is discovered in later life.

The classification adopted by the New York Heart Association in 1924 whereby heart disease in pregnancy is divided in to four grades has proved helpful. Grade 1 is symptomless grade 2 is subdivided into those with slight symptoms (grade 2a) and those with more severe symptoms but not yet having congestive failure or auricular fibrillation (grade 2b) grade 3 signifies auricular fibrillation or heart failure past or present. No restrictions are placed on women in grade 1. Women in grade 3 are strongly dissuaded from pregnancy and termination is usually advised if they are already pregnant. Women in grade 2 may present more awkward problems.

The chief difficulty arises out of the instability of the grading. A woman starting pregnancy in grade 2a usually ends it in grade 2b. If she starts in grade 2b she may later enter grade 3. All these cases must be carefully watched. Any tendency towards increase of symptoms is met by increased rest even to complete rest in bed. In this way the great majority can be kept in their original grade. A woman in grade 2a should not be dissuaded from pregnancy but may be advised to limit her family to two or three.

Women in grade 2b are usually warned against pregnancy. If they have already conceived pregnancy should be terminated during the first three months. If gestation is already between the third and sixth month they should be put to bed and observed for a week or two. If they improve rapidly and enter grade 2a pregnancy is allowed to continue. If they do not improve pregnancy should be terminated. It must be borne in mind that conditions steadily deteriorate up to the end of the sixth month so that a woman in grade 2b at the end of the fourth month failing to respond adequately to sufficient rest is almost certain to enter grade 3 in the sixth month. Management of pregnancy after the sixth month when the patient is in grade 2b depends upon the history of the pregnancy and the response to rest. If seen for the first time and if previously physically active such a woman will surely enter grade 2a with rest in bed and she may be taken safely to term. But if already under observation and receiving adequate rest further improvement cannot be expected and the continued strain of the next three months may well induce auricular fibrillation or heart failure. She should remain strictly in bed. If she improves she may be taken to term. If she remains problematical it is wise to terminate at the end of the eighth month. If she deteriorates pregnancy should be terminated as soon as heart failure or auricular fibrillation are controlled.

In a series of 546 cases of heart disease in pregnancy of which 472 were rheumatic the respective mortality rates for class 3 2b 2a and 1 were 40 per cent 4.7 per cent 0.5 per cent and nil (Pardee 1934). Hamilton (1947) reported more or less similar figures in a series of 1335 cases of which 93 per cent were rheumatic. When the cardiac findings were favourable (grade 1 and 2a) the mortality rate was 2 per cent and was the same

as in non pregnant controls in the same category when the cardiac findings were unfavourable (grades 2b and 3) the mortality was 18 per cent compared with 6.7 per cent in non pregnant controls. The infant mortality was 8.6 per cent in favourable cases and 31 per cent in the unfavourable. When there was auricular fibrillation the maternal mortality was 32 per cent (8 per cent in controls) and the infant mortality was 50 per cent. The most common causes of death in the unfavourable group were congestive heart failure (64 per cent) and embolism (13 per cent). Hamilton also noted that the mortality rate had increased slightly (from 16 to 18 per cent) since termination in the last trimester had been given up. No difference in prognosis between cases of mitral stenosis and aortic incompetence was noted in any of the series mentioned.

The risk of pregnancy in rheumatic cases does not end with the birth of the infant. Pulmonary embolism in particular is more likely to occur during the puerperium.

Cases of active rheumatic carditis are probably best terminated as soon as the state of the heart permits for there is no knowing what the subsequent course will be and a relapse later in pregnancy may prove very serious.

When pregnancy is not advised prevention is best insured by a simple sterilising operation. Termination of pregnancy is by therapeutic abortion in the first three months by abdominal hysterotomy from the fourth to the sixth month by induced labour or by Cæsarean section during the seventh and eighth months by natural means or by Cæsarean section at term. The choice must rest with the obstetrician.

BACTERIAL ENDOCARDITIS

Before the introduction of penicillin the life of the fetus was the main consideration. The situation is now reversed however and every effort should be made to save the mother. As heart failure is now the chief cause of death from bacterial endocarditis termination of pregnancy may often be desirable.

THYROTOXICOSIS

One of the few known factors that may aggravate or precipitate thyrotoxicosis is pregnancy. It follows that thyrotoxic women should be advised against pregnancy until they are cured. Improvement on rest and iodine or as a result of thiouracil treatment is not enough such cases tend to relapse during pregnancy. At least a year should pass after partial thyroidectomy or thiouracil cure before conception should be considered.

If a woman is thyrotoxic and already pregnant therapeutic abortion should be considered during the first three months if not seen until gestation is more advanced it may be wiser to take the patient to term with the aid of thiouracil. Subtotal thyroidectomy is better deferred owing to the risk of relapse. The dose of thiouracil must be the minimum that is effec

tive for there is some danger of its causing goitre in the foetus the simultaneous administration of small doses of iodine or thyroid may prevent this (see page 493)

HYPERTENSION

High blood pressure discovered during pregnancy may be due to chronic persistent hypertension (usually essential) or to toxæmia of pregnancy. Essential hypertension may be aggravated by pregnancy but with rest diet and sedatives mild cases can be taken to term. Nevertheless women with high basal blood pressures (above 160/100 mm Hg) should be advised against pregnancy in view of the increased risk of toxæmia the high infant mortality (66 per cent according to Browne 1947) and the chances of serious aggravation. For similar reasons pregnancy should be terminated in women with relatively high pressures in the first three months. Hypertension associated with toxæmia of pregnancy is a separate problem and will not be considered here.

ARTERIO VENOUS ANEURYSM

Arterio venous aneurysm may be congenital (cirroid) or acquired (usually as a result of a perforating wound) and may occur in any situation particularly in the brain limbs or lung.

CONGENITAL CIRROID ANEURYSM

Cirroid aneurysm consists of a twisted mass of vessels in which arteries and veins are in direct communication. One or more superficial hæmangiomas may be seen elsewhere or there may be a family history of such nævi.

The cerebral type may give rise to epilepsy to subarachnoid hæmorrhage or to ophthalmoplegic migraine. Examination may reveal a systolic murmur heard best through the eye ball on the affected side or sometimes over the skull. The diagnosis may be proved by finding an unduly high oxygen saturation in samples of blood obtained from the ipsilateral jugular vein. The lesion may be localised by means of angiography 10 to 20 ml of 70 per cent diodone or other radio opaque substance being injected rapidly into the carotid artery and skiagrams of the cerebral vessels being obtained at the appropriate moment. The condition should be distinguished from berry aneurysm and from Sturge's disease in which facial and pial nævi without arterio venous communications are associated with calcification of brain substance epilepsy mental retardation and glaucoma (Nussey and Miller 1939). Treatment consists of ligation of the common carotid artery on the side of the lesion if after trial compression hemiplegia or other serious ischæmic symptoms do not occur. The risk of such an untoward event increases progressively with the age of the patient.

Cirroid aneurysm in a limb presents similar features to those of its trau



Fig. 19.05 (a)—Diagram showing a congenital arterio venous aneurysm of the lung. The appearances bear some resemblance to those of pulmonary tuberculosis
(b) Angiocardiogram showing diodone filling the aneurysm



Fig. 19.06—Calcification in the wall of an arterio venous aneurysm

matic cousin. It may be situated anywhere from the shoulder or pelvic girdle to the hand or foot. There is usually an increase in blood flow to the limb which may be longer and larger than its fellow. The veins stand out and may pulsate and the skin temperature is raised. It may be possible to locate the aneurysm with precision by observing the effect on the local and general circulation of compressing the various arteries of the limb at appropriate points. An impressive machinery murmur and thrill may be appreciated over the aneurysm itself. Venous blood from the affected limb may be more saturated with oxygen than venous blood from the unaffected limb. The exact location and construction of the aneurysm may be demonstrated by means of angiography. Treatment is more difficult than in traumatic cases. Excision is usually impossible owing to the diffuse nature of the lesion; moreover affected vessels are physiologically abnormal and fail to constrict when injured so that severe and prolonged hæmorrhage may follow surgical interference. Ligation of the main vessels leading to the aneurysm (above and below) may be possible but deep X-ray therapy is usually best.

Congenital arterio venous aneurysm in the lung causes venous blood from the pulmonary artery to be shunted directly into the pulmonary veins and hence into the arterial circulation at the same time the blood flow through the rest of lung may be reduced the steep pressure gradient through the aneurysm offering the easier pathway. The result is a lowered arterial oxygen saturation in the region of 70 to 75 per cent (Burchell and Clagett 1947) central cyanosis polycythæmia and clubbing. Most of the cases reported have been in children or young adults. Hæmoptysis has occurred in 50 per cent. The heart itself is normal but there is often a continuous machinery murmur over the affected part of the lung. A skiagram may show a shadow not unlike local chronic pulmonary tuberculosis (fig 19.05a) but on fluoroscopy the lesion may be seen to pulsate and angiocardiograms may show the abnormal vessels filled with diiodone (fig 19.05b). Lesions may be single or multiple and unilateral or bilateral. Calcification may occur in the wall of an aneurysm (fig 19.06). One case (a girl aged 9) seen by the author died with cerebral abscess (see page 243). The condition should be distinguished from patent ductus arteriosus helping to correct pulmonary or tricuspid atresia. Treatment by lobectomy or pneumonectomy is curative unless there are several widely distributed aneurysms (Barnes *et al.* 1948).

ACQUIRED ARTERIO VENOUS ANEURYSM

The great majority of acquired arterio venous aneurysms are due to perforating gunshot wounds in war and are seen most often in connexion with the femoral brachial or carotid arteries. Occasionally they may be syphilitic mycotic or artificial. Arterio venous shunting may also occur in highly vascular structures such as the thyroid gland in severe thyrotoxicosis or as a result of overdosage with thiouracil (page 493) the uterus in

pregnancy (page 507) and the bones in active Paget's disease (page 517)

The local signs and the effect on the general circulation are similar to those in congenital circoid aneurysm but will be described more fully here because most investigations on the effects of arterio venous shunting have been carried out on traumatic cases usually with lesions in the thigh

Local signs in the affected limb include fullness of the veins increase of skin temperature and sometimes œdema. On the other hand, peripheral ischaemic symptoms may predominate and the toes may be unduly cold or even gangrenous (in cases of recent origin). A gross machinery murmur and thrill are invariable over the aneurysm. The variability of the blood flow in the affected limb as judged by clinical criteria has been confirmed by Cohen Edholm and others (1948) who found reduced flows distal to the lesion in two relatively recent cases and an increased flow in a case of 29 years' duration. The blood flow in the unaffected limbs was normal.

The general circulation is hyperkinetic (page 502) and if the shunt is large enough paroxysmal cardiac dyspnoea or signs of congestive heart failure may develop. If the shunt is temporarily obliterated by digital compression of the femoral artery just above the lesion the pulse rate falls 10 to 20 beats per minute (Branham's sign) the blood pressure rises 10 to 15 mm Hg the venous pressure falls slightly and the cardiac output falls (Stead and Warren 1945) but capillary pulsation is accentuated (Lewis and Drury 1923). Slowing of the pulse may be due to the inhibiting effect of the abrupt although slight fall in right auricular pressure on the Bainbridge reflex and is said to be abolished by 2 to 3 mg of atropine (Kramer and Kahn 1946).

Cardiac enlargement is almost certainly due to the raised cardiac output and increased stroke volume. The total blood volume is also increased in many cases (Holman 1937). The hyperkinetic circulation is maintained by tachycardia and raised venous filling pressure whilst the peripheral resistance is reduced by vasodilatation in skin and muscle.

Treatment. Any arterio venous aneurysm large enough to influence the general circulation should be repaired. Smaller lesions may be left alone if causing no local symptoms and some of them become obliterated spontaneously. Every effort should be made to repair the artery by lateral suture with or without the aid of a covering strip of vein so that the normal circulation is preserved (Junghanns 1943). Ligation of artery and vein above and below the aneurysm is less satisfactory the resulting circulation through the brain or limb being sometimes inadequate.

THE HEART AND CIRCULATION IN BERI BERI

In modern civilised communities pure beri beri is rare the clinical picture being commonly influenced by deficiencies in vitamins other than aneurin (B_1) and by associated conditions especially chronic alcoholism.

Aneurin (thiamine) in association with other components of the vitamin B complex is found chiefly in unpolished rice Marmite liver yeast wheat and other grains It is used by the body in carbohydrate metabolism its chief known function being concerned with the oxidation of pyruvic acid which is formed from lactate When there is insufficient aneurin carbohydrate metabolism is held up at this point and an excess of pyruvic acid accumulates in the blood (Peters 1939) It follows that any condition in which carbohydrate metabolism is excessive predisposes to beri beri in that aneurin requirements are heavier When in addition the vitamin B intake is reduced at the same time as in chronic alcoholism vomiting of pregnancy and thyrotoxic crises beri beri may well develop

The normal requirement of aneurin is about 1 mg daily for an adult and is supplied adequately by the ordinary European diet Special ulcer diets however, unless supplemented may be deficient and psychoneurotic patients with severe anorexia and vomiting may not receive a sufficient supply of the vitamin Beri beri was common in German concentration camps and Japanese prison camps during the second world war although usually complicated by other vitamin deficiencies and has always been relatively common in the Far East when the basic food has been polished rice

Aneurin deficiency is rarely gross in civilised communities and so the presence of some additional factor is commonly needed before the effects of slight deficiencies are brought to light Under these conditions beri beri is atypical for such patients are apt to be middle aged or elderly and the classical signs may be masked by hypertension coronary sclerosis or emphysema Again although beri beri especially affects the right side of the heart in pure cases it will affect the left side if left sided stress is already present as in hypertension or if the blood supply to the left ventricle is deficient as in occlusive coronary atherosclerosis Thus in these mixed cases no clear picture of beri beri develops (Konstam and Sinclair 1940)

Behaviour of the heart and circulation The pure disease was studied in Java by Wenckebach (1928 1934) The essential features included a hyperkinetic circulation vasodilatation enlargement of the right side of the heart and dilatation of the pulmonary artery Few accurate cardiac output studies have been carried out but the clinical description and the swift circulation time (Weiss and Wilkins 1936-37) leave little doubt that it is high Heart failure may develop suddenly and fulminating cases occur in which death results within 24 to 48 hours of the alleged onset of symptoms (Hashimoto 1937) Even in Great Britain cases have been described in which heart failure has occurred remarkably suddenly and unexpectedly leading to a rapidly fatal issue (Wood 1939)

The cause of the hyperkinetic circulation is not yet clear It may result from the same unidentified mechanism that ensures a high cardiac output to compensate for tissue hypoxia in anæmia and anoxic cor pulmonale but instead of hypoxia there is faulty carbohydrate metabolism and the high

cardiac output therefore achieves nothing. The drop in venous pressure and quietening of the circulation that follow the injection of 1 ml of pitressin and the stormy reaction to 1 mg of subcutaneous adrenaline (Wenckebach 1928) suggest that vasodilatation in muscle may be directly responsible by causing an arterio-venous shunt. That some such mechanism exists is suggested by the sudden rise in pulse rate, venous pressure and cardiac output that may follow the subcutaneous injection of 10 mg of mecholin which is known to cause vasodilatation in muscle.

The heart itself shows little specific at necropsy, the disturbance being biochemical not structural.

Diagnosis. The clinical diagnosis of cardiovascular beri beri rests on an appropriate dietetic history, the demonstration of a hyperkinetic circulation, radiological appearances showing conspicuous dilatation of the pulmonary artery and right ventricle, electrocardiographic evidence of right ventricular stress, the response to pitressin and adrenaline, associated polyneuritis and on the finding of a raised blood pyruvic acid or reduced amounts of aneurin in blood (Jansen 1938, Sinclair 1938) or urine (Harris *et al.* 1938, McAlpine and Hills 1941).

Peripheral neuritis usually begins with pain in the calves on walking, similar in character to intermittent claudication. Associated weakness of the legs, marked tenderness of the calves, numbness and tingling of the fingers and toes, loss of deep tendon jerks and glove and stocking anaesthesia are usually found.

Evidence of deficiencies in other vitamins, especially of the vitamin B group, is helpful in proving inadequacy of the diet.

Treatment. It must be stressed that the symptoms of beri beri may begin abruptly and that the course of the disease may be fulminating, death occurring within a few days of the onset. Once the diagnosis has been made there may be no time to lose. Again, the possibility of vitamin B₁ deficiency should always be borne in mind in any case of heart failure of obscure origin, especially when right-sided. Here is one of the fatal forms of heart disease which is curable.

The patient should be put to bed immediately and aneurine hydrochloride should be given at once intravenously in an initial dose of 50 to 100 mg. The effect is dramatic if not given too late. Subsequent doses should be of the order of 10 to 20 mg. per day for a fortnight orally or parenterally and followed by an adequate diet. An abundance of the other components of the vitamin B group is also advised.

Fulminating cases should benefit by repeated injections of pitressin (1 ml. 4 hourly) until the vitamin has had time to work, but care must be taken to avoid hydraemia by keeping the salt and water intake as low as possible.

Chronic alcoholics, cases of severe thyrotoxicosis, Simmond's disease or anorexic nervosa and women vomiting in pregnancy should be given 2 to 3 mg. of aneurin daily as a precautionary measure.

PAGET'S DISEASE OF BONE

The hyperkinetic circulation associated with extensive active Paget disease was first clearly demonstrated by Edholm, Howarth and McMichael in 1945. The general cardiovascular findings closely simulate those associated with arterio-venous aneurysm. In the case described by Edholm *et al.* the blood flow through actively diseased bones was estimated to be 3 to 4 litres per minute and the total cardiac output was 13 litres per minute. The venous pressure was elevated and there was dependent oedema. Further observations on other cases of active Paget's disease have shown that the heart is not usually overloaded for it is capable of increasing its output by means of tachycardia or a greater rise of venous filling pressure. On the other hand, paroxysmal cardiac dyspnoea may then occur (McMichael 1947).

Paget's disease also encourages metastatic calcification, especially Monckeberg's sclerosis and calcification of the valve rings of the heart. Extension to the interventricular septum may involve the bundle of His or its branches with the production of complete heart block or bundle branch block respectively (Harrison and Lennox 1948).

Cor pulmonale secondary to thoracic deformity from Paget's disease has also been described (Wilks 1869).

Diagnosis. If aortic incompetence and valve calcification are both present the clinical diagnosis of Paget's disease may be overlooked in favour of atherosclerotic aortic valve disease. As long as the condition is borne in mind, however, diagnosis is easy for skiagrams of the bones show characteristic changes and the blood alkaline phosphatase is very high.

HEPATIC FAILURE

It is becoming increasingly evident that advanced disease of the liver may lead to a hyperkinetic circulatory state in addition to the well known palmar flush and cutaneous spider naevi. The usual cause is secondary carcinoma but common cirrhosis and even serious infective hepatitis may be responsible. It appears that the liver normally detoxicates some vaso-depressor substance and that this substance accumulates when the organ is failing. Vasodilatation results in certain territories such as skin and muscle and it is likely that arterio-venous communications open up and cause an extensive arterio-venous shunt. If unrecognised the situation may lead to embarrassing diagnostic error, the raised venous pressure, oedema and enlargement of the liver being readily attributed to heart failure.

REFERENCES

THE HEART IN ANÆMIA

Block C (1937) Heart involvement and electrocardiographic findings in anæmia *Acta med Scand* 93 543

Bouchut L and Frément R (1934) Les gros cœurs peu anoxémiques à propos des anémies pernicieuses compliquées d'hypertrophie et d'insuffisance cardiaques *Arch Mal du Cœur* 27 325

Cabot R C and Richardson O (1919) Cardiac hypertrophy in pernicious anæmia *J Amer med Ass* 72 991

Coombs C F (1926) The cardiac symptoms of pernicious anæmia with particular reference to cardiac pain *Brit med J* ii 185

Forman M B and Daniels A L (1930-1) Effect of nutritional anæmia on size of the heart *Proc Soc exper Biol and Med* 28 479

Goldstein B and Boas E P (1927) Functional diastolic murmurs and cardiac enlargement in severe anæmias *Arch intern Med* 39 226

Grunberg F W (1930) Über einige Veränderungen von seiten des Herzgefäßsystems bei Schweren anämien *Deutsch Arch f klin Med* 169 354

Harrison T R (1935) Failure of the circulation Baltimore

Kraus F (1905) Die klinische Bedeutung der fettigen Degeneration des Herzmuskels Schwer anämischer Individuen *Berl klin Wchnschr* 42 5

Liljestrand G and Stenstrom N (1925-6) Work of heart during rest influence of variations in hæmoglobin content of blood flow *Acta med Scand* 63 130

McMichael J (1947) Circulatory failure studies by means of venous catheterization *Advances in Internal Medicine* 2 64 — Sharpey Schafer E P Mollison P L and Vaughan J M (1943) Blood volume in chronic anæmia *Lancet* i 637

Nielson H E (1934) The circulation in anæmic conditions *Acta med Scand* 81 571

Pickering G W and Wayne E J (1934) Observations on angina pectoris and intermittent claudication in anæmia *Heart* i 3

Sahl H (1895) Ueber diastolische accidentelle Herzgeräusche *Blatt f Schweizer Aerzte* 25 33

Sharpey Schafer E P (1944) Cardiac output in severe anæmia *Chn Sc* 5 125

Smith K S (1933) Nutrition of heart in relation to electrocardiogram and anginal pain *Lancet* i 632

von Noorden C (1891) Untersuchungen über Schwere Anämien *Charité Annalen* 16 217

THE HEART IN PREGNANCY

Burns J J and Rubricus J (1948) The determination of the prognosis of pregnancy in rheumatic heart disease *Amer Heart J* 35 28

Browne F J (1947) Chronic hypertension in pregnancy *Brit med J* ii 283

Burwell C S Strayhorn W D Flickinger Corlette M B Baverman E P and Kennedy J A (1938) Circulation during pregnancy *Arch intern Med* 62 979

Hamilton B E (1947) Report from the cardiac clinic of the Boston lying in hospital for the first twenty five years *Amer Heart J* 33 663

Jensen J (1938) The heart in pregnancy London

Pardee H E B (1934) Cardiac conditions indicating therapeutic abortion
J Amer med Ass 103 1899

ARTERIOVENOUS ANEURYSM

Barnes C G Fattu I and Pryce D M (1948) Arteriovenous aneurysm of the lung *Thorax* 3 148

Burchell H B and Clagett O T (1947) The clinical syndrome associated with pulmonary arteriovenous fistulas including a case report of a surgical cure
Amer Heart J 34 151

Cohen S M Edholm O G Howarth S McMichael J and Sharpev Schafer E P (1948) Cardiac output and peripheral blood flow in arteriovenous aneurysm *Clin Sc* 7 35

Holman E (1937) Arteriovenous aneurysm New York

Junghanns H (1943) Lateral suture in carotid aneurysm after gunshot wound *Arch f klin chirurg* 205 149

Kramer M L and Kahn J W (1946) Effect of atropine on the Branham sign in arteriovenous fistula *Arch intern Med* 87 8

Lewis T and Drury A V (1923) Observations on arterio venous aneurysm
Heart 10 307

Nussev A M and Miller H H (1939) Sturge's disease *Brit med J* 1 8--

Stead E A and Warren J V (1945) Circulation before and after operation for arteriovenous fistula Committee on Med Research Bull New York 64 711

THE HEART AND CIRCULATION IN BERI BERI

Harris L J Leong P C and Ungley C C (1938) Measurement of vitamin B₁ in human urine as an index of the nutritional level *Lancet* 1 539

Hashimoto H (1937) Acute pernicious form of beri beri and its treatment by intravenous administration of vitamin B with special reference to electrocardiographic changes *Amer Heart J* 13 580

Jansen B C P (1938) Chemical determination of aneurin (vitamin B₁) in blood *Acta brev Neerland* 8 119

Konstam G and Sinclair H M (1940) Cardiovascular disturbances caused by deficiency of vitamin B *Brit Heart J* 2 231

McAlpine D and Hills G M (1941) The clinical value of the thiochrome test for aneurin (vitamin B) in the urine *Quart J Med* 10 31

Peters R A (1939) Discussion on the clinical aspects of the vitamin B complex *Proc Roy Soc Med* 32 807

Sinclair H M (1938) Value of estimation of vitamin B in blood *Quart J Med* 7 591

Weiss S and Wilkins R W (1936) The nature of the cardiovascular disturbances in vitamin deficiency state *Trans Ass Amer Phys* 2 341 ——— (1937) Disturbance of the cardiovascular system in nutritional deficiency *J Amer med Ass* 109 786 ——— (1937) The nature of the cardiovascular disturbances in nutritional deficiency states (beri beri) *Ann intern Med* 2 104

Wenckebach K F (1928) St Cyres lecture on heart and circulation in tropical avitaminosis (beri beri) *Lancet* ii 265 ——— (1934) Das Beriberi Herz Berlin

Wood P H (1939) The effect of vitamin B deficiency upon the cardiovascular system *Proc Roy Soc Med* 32 817

PAGET'S DISEASE OF BONE

Edholm O G, Howarth S and McMichael J (1943) Heart failure and bone blood flow in osteitis deformans *Chin Sr* 5 249

Harrison C V and Lennox B (1948) Heart block in osteitis deformans *Brit Heart J* 10 167

McMichael J (1947) Circulatory failure studies by means of venous catheterisation *Advances in Internal Medicine* 2 64

Wilks S (1869) Case of osteoporosis or spongy hypertrophy of the bones (calvaria clavicle os femoris and rib exhibited at the society) *Trans path Soc of London* 20 273

TRAUMATIC LESIONS OF THE HEART AND
GREAT VESSELS

SPONTANEOUS LESIONS

SPONTANEOUS traumatic lesions of the heart or great vessels include dissecting aneurysm of the aorta rupture of a hypoplastic aorta or syphilitic aortic aneurysm ruptured valve cusps in bacterial endocarditis rupture of a congenital syphilitic or mycotic aneurysm of a sinus of Valsalva into the right side of the heart rupture of chordae tendineae in rheumatic or bacterial endocarditis and rupture or perforation of the heart or ventricular septum secondary to cardiac infarction or ventricular aneurysm. The majority of such lesions have been described elsewhere as complications of the diseases mentioned. Only dissecting aneurysm and rupture of an aneurysm of a sinus of Valsalva into the right side of the heart remain to be considered here.

DISSECTING ANEURYSM

Definition Dissecting aneurysm was so called by Lænnec (1826) and means dissection of the media of the aorta by extravasated blood that has penetrated between its coats from the vasa vasorum or from the lumen of the vessel.

Incidence About 1 per cent of all sudden deaths are due to dissecting aneurysm (Mote and Carr 1942). Hospital records which include relatively few such deaths give an approximate incidence of one dissecting aneurysm in every 450 necropsies. Men are more susceptible than women in the ratio of 3 : 2. Patients are commonly between 50 and 60 years old but 24 per cent are under 40 (Schnikter and Bayer 1944) and a case has been recorded in a boy of 15 (Galbraith Gardner and Hardwick 1939). About 50 per cent of dissecting aneurysms in women have occurred during pregnancy (Schnikter and Bayer 1944).

Etiology and pathology Virchow's original conception that dissection follows an intimal tear at the site of an atheromatous ulcer is no longer tenable for a tear at such a site is now known to be rare (Shennan 1934). Although hypertension and atheroma are usually associated they are not essential the intima may be normal and not even ruptured (Tyson 1931).

Dissection is always within the media commonly begins in the ascending aorta and appears to be closely related to cystic medial necrosis (Erdheim 1929). The cause of such necrosis is unknown. Tyson's thesis that it was due to obliterative endarteritis of the vasa vasorum has not been con-

firmed Cystic necrosis without dissection may be found sometimes in routine necropsies (Moritz 1932 Rottino 1939) Whether hæmorrhage into the diseased media commonly follows an intimal tear, or whether it comes from the vasa vasorum (the intimal tear then being due to secondary rupture) remains uncertain When the intima is intact hæmorrhage obviously cannot come from the lumen of the aorta On the other hand intimal tears may undoubtedly be primary for they may occur in healthy ascending aortas without subsequent dissection (Peery 1942) Occasionally hæmorrhage occurs into an area of cystic necrosis of the media without dissection the hæmatoma then becoming organised and causing no trouble (Shennan 1934)

Dissection may spread proximally and involve the root of the aorta causing aortic incompetence occasionally the coronary arteries are dissected and occluded Dissection usually spreads distally however may travel the whole length of the aorta and may proceed along any of its branches Ischæmic effects from occluded visceral or parietal vessels are common The majority of cases die from external rupture usually into the pericardium sometimes into the left pleural cavity or elsewhere Occasionally dissection associated with an intimal tear in the ascending aorta ruptures back into the lumen of the vessel at some distal point forming an alternative or double aortic channel (double barrelled aorta) This is found in the majority of cases which recover (Shennan 1934)

Clinical features Dissection of the aorta may be precipitated by effort (Gager 1928) and give rise to a well defined clinical picture with characteristic variations A typical attack begins suddenly with severe pain in the centre of the chest or in the præcordial area The pain may be gripping tearing shooting or vice like and usually lasts for hours it may radiate to the head and neck to the back—less often to the arms Later in the attack it may spread to the lumbar regions or abdomen and occasionally into the legs depending on the extent of the dissection In perhaps half the cases however pain is slight or absent (Baer and Goldburgh 1948)

Breathlessness is nearly as common as pain and syncope is not rare (Hamburger and Ferris 1938) Attacks may therefore closely resemble coronary thrombosis but in cases which survive the blood pressure usually remains high and the electrocardiogram normal moreover dilatation of the aorta may often be seen in skiagrams (Wood Pendergrass and Ostrum 1935)

Other findings depend upon the site and extent of the dissection upon which branches of the aorta are occluded and upon the site of external rupture Aortic incompetence may develop when the root of the aorta is dissected (Weiss 1935) and is being noted with increasing frequency (David *et al* 1947) myocardial infarction may occur if the left or right coronary artery is occluded giving rise to the appropriate electrocardiographic pattern (Wainwright 1944) Pericardial friction is heard occasionally and hæmopericardium may be recognised before death

Dissection of major arteries leads either to occlusion of the vessel or to

increased amplitude of pulsation due to spontaneous periarterial sympathectomy (Weisman and Adams 1944) Occlusion of one or other or both carotid arteries may cause hemiplegia mental confusion or coma of the anterior spinal artery, paraplegia of arteries to the limbs loss of the peripheral pulse and perhaps ischæmic pain of the renal artery hæmaturia—and so on Occasionally a pulse that has been absent may re appear as a result of rupture re entry (Lawrence 1935) A systolic murmur and thrill may develop over partly occluded vessels including the aorta (McGeachy and Paullin 1937) Left hæmothorax is found in about 12 per cent of cases (Baer and Goldburgh 1948) Hæmorrhage into the mediastinum may be responsible for cough and dysphagia An abdominal mass may become palpable Hæmoptysis hæmatemesis and hæmaturia occur occasionally

Cases which survive the original dissection may present themselves later with congestive heart failure associated with aortic incompetence When there has been no history of pain such cases have usually been diagnosed erroneously as syphilitic aortic incompetence despite negative Wassermann reactions (Gouley and Anderson 1940 Flaxman 1942)

Finally in differential diagnosis it should be remembered that fever and leucocytosis are the rule during the first few days not the exception (Baer and Goldburgh 1948)

Prognosis According to Shennan (1934) about 10 per cent of all cases of dissecting aneurysm recover from the attack usually owing to rupture re entry The majority succumb later to heart failure either as a result of aortic incompetence or from associated hypertensive heart disease

Treatment No treatment is likely to influence the course of dissection Morphine should be given freely to combat pain If the patient survives the initial attack he should be kept in bed for at least three weeks

RUPTURE OF AN ANEURYSM OF AN AORTIC SINUS (SINUS OF VALSALVA) INTO THE RIGHT VENTRICLE RIGHT AURICLE OR PULMONARY ARTERY

Aneurysm of one of the aortic sinuses may be congenital syphilitic or mycotic Rupture of such an aneurysm into the pericardium or left pleural cavity is immediately fatal but perforation into the right auricle ventricle or pulmonary artery leads to a well defined clinical syndrome which may be compatible with many years of active life

Incidence The condition is rare indeed the author has only encountered and investigated four living instances (Congenital cases may occur in young adults syphilitic cases in later life and mycotic at any age)

Physiology Rupture into the right auricle causes a high pressure in that chamber overloading of the right heart and the rapid development of congestive failure Samples of blood obtained by means of cardiac catheterisation should be similar to those obtained in atrial septal defect (page 218) Perforation into the right ventricle may similarly overload the right



(a) Anteroposterior view showing engorged pulmonary circulation enlargement of the left ventricle and resection of the 5th rib on the left side (the case having been operated on for patent ductus)



(b) Second oblique view showing enlargement of the left ventricle and dilatation of the pulmonary artery

Fig. 20.01—Case of ruptured mycotic aneurysm of aortic sinus into the pulmonary artery

heart blood samples and intracardiac pressures are similar to those in ventricular septal defect (RAP 0 RVP 17 PAP 21 cm saline SVC and RA samples 44 to 45 RV and PA samples 28 ml O₂ unsat per litre in a case seen by the author)

Perforation into the pulmonary artery sets up similar features to patent ductus arteriosus (fig. 20.01). In one such case investigated by the author due to a perforated mycotic aneurysm from bacterial endocarditis (cured by penicillin) samples from the right auricle and ventricle showed 67 to 70 ml oxygen unsaturation per litre whereas pulmonary artery samples were only 33 to 36 ml unsaturated. The mean right ventricular pressure was 40 to 43 cm of saline above the sternal angle and the pulmonary artery pressure plus 86 cm.

Clinical features Pain may occur from involvement of the orifice of one or other coronary arteries but is otherwise absent. The onset is usually signalled by the rapid development of congestive heart failure but not necessarily. The two cases mentioned above were by no means incapacitated and had both lived seven years since the onset.

The chief signs are a loud machinery murmur accompanied by a thrill over the base of the heart but at a lower level than that associated with

patent ductus arteriosus accompanied by signs of aortic incompetence and by features resembling those of ventricular septal defect or patent ductus according to the site of the perforation

Prognosis Rapid deterioration to a fatal outcome is said to be the rule (Abbott 1919) but this may be because the diagnosis is usually only made at autopsy. The author's four cases are not only alive but relatively well

EFFICACY OF DIRECT INJURY

Direct injury to the heart may be caused by stab or gunshot wounds and very rarely by diagnostic procedures such as needling the pericardium. The literature on the subject has been well surveyed by King (1941) and by Barber (1944)

GUNSHOT WOUNDS

A bullet or piece of shrapnel may perforate the heart through and through, may lodge in the myocardium or pericardium with or without perforation of one or more chambers, or may graze the surface of the heart without causing death. In an analysis of 25 instances of war wounds involving the heart made in conjunction with Nicholson in 1945 the relative incidence of such lesions was as follows

Near misses	4
Grazes or tangential wounds	4
Through and through perforation	3
Foreign body in pericardium	7
Foreign body in myocardium	7

Of 1640 consecutive penetrating chest wounds treated at Nicholson's centre the heart was directly or indirectly injured in 17 per cent. The immediate result is hæmopericardium and the rapid development of cardiac tamponade. If a foreign body passes close to the heart or lodges within half an inch of its surface a transient pericardial serous effusion may develop. If the patient does not die from cardiac tamponade or hæmorrhage into the pleural cavity complete recovery may follow whether or not a metallic foreign body remains in the heart.

The chief complication during convalescence is recurrent acute pericarditis; this is nearly always associated with the presence of a foreign body either in the pericardium or closely connected with it (Wood 1945); it rarely arises when a bullet is embedded deeply in the myocardium. The attacks tend to be severe, with pain, fever, tachycardia, gross electrocardiographic changes, and the rapid development of a sterile serous effusion which may cause cardiac tamponade. They usually last about a week. The first attack may occur at any time during convalescence up to about three months after the injury and may recur several times at intervals of about a month. Of five such cases studied by the author in the second world war

all finally recovered, three without interference and two after removal of the foreign body by Nicholson (1945)

A second complication is coronary thrombosis during convalescence when a pericardial foreign body is lodged in contact with a major coronary vessel but this was observed only once

Intravascular foreign body has been described on page 438

Diagnosis The possibility of cardiac injury should be considered in all cases of gunshot wounds of the trunk or neck especially if the missile is judged to have been directed towards the heart or if its direction is not known for certain. Early diagnosis depends upon recognising the signs of cardiac tamponade or hæmopericardium (page 343). An electrocardiogram may be most helpful by showing the presence or absence of the pericardial T_w pattern

Intracardiac or pericardial foreign body may be readily detected by means of fluoroscopy but may be easily overlooked in skiagrams

Treatment It is impossible to say how many lives might be saved by early surgical repair of cardiac wounds. In the second world war the majority of recognised cases survived without such early repair or at least lived long enough to be evacuated to general hospitals. They were therefore

all relatively favourable cases and the great majority recovered

Relief of cardiac tamponade by paracentesis may be life saving both in the early stages or during a later attack of acute pericarditis. Metallic foreign bodies lodged in the pericardium are best removed in view of the danger of recurrent pericarditis. Although none of the attacks witnessed proved fatal the episodes were most alarming. Intracardiac foreign bodies should probably be removed if superficial and left alone if deep

Prognosis Only one of the twenty five patients mentioned previously died but as already stated these were favourable cases in that they had survived



Fig. 200 —Skiagram showing machine gun bullet embedded in the wall of the right auricle

until evacuated to a general hospital

Follow up studies are incomplete but the worst case with three attacks of recurrent pericarditis and a machine gun bullet embedded in the wall

of the right auricle (fig 20 02) was alive and well two years after being wounded

In 1937 the author had the opportunity of investigating a healthy man with a machine gun bullet embedded in his heart since 1917 (fig 20 03) This case was reported in detail by Grey Turner (1941) On the whole it seems likely that the ultimate fate of these patients is favourable



Fig 20 03—Skigram showing machine gun bullet embedded in the heart since 1917 (see text)

STAB WOUNDS OF THE HEART

Direct injury to the heart in civil life is usually due to single or multiple stab wounds the majority of which penetrate the right ventricle The clinical physiological radiological and electrocardiographic features of cases which have survived long enough to receive medical aid have been chiefly those of hæmopericardium (Wood 1937) Death from hæmorrhage into the pleural cavity or from cardiac tamponade may be prevented by timely surgical repair

Even when patients appear to be holding their own it is probably wise to evacuate the blood clot and to repair and sterilise the wound as soon as possible, for hæmorrhage may continue or recur and serious cardiac tam-

ponade develops in most cases Moreover if tamponade is unrelieved too long acute coronary insufficiency may seriously impair the function of the myocardium and when it is finally relieved death may result from acute heart failure The development of a bulge on the left border of the heart simulating the appearances of ventricular aneurysm should not deter the surgeon for this is likely to prove no more than a localised pericardial hæmatoma (fig 20 04)



Fig 20 04—Localised pericardial hæmatoma superficially resembling a cardiac aneurysm

EFFECTS OF INDIRECT INJURY

Indirect injury to the heart may be caused by crushes blows falls or blast. The effects include sudden death from ventricular fibrillation or standstill rupture of the aorta, rupture of one or more chambers of the heart rupture of the aortic or mitral valve hæmopericardium, myocardial bruising, auricular fibrillation and heart block. Coronary occlusion and subsequent angina pectoris or cardiac infarction may also occur but the relationship to trauma is less well understood.

SUDDEN DEATH

A heavy blow to the region covering the heart may cause sudden death from ventricular fibrillation or cardiac rupture both naturally and experimentally in dogs (Bright and Beck 1935).

There have been numerous instances of sudden death resulting from relatively minor trauma of a kind quite incapable of damaging the heart. The catastrophe is then ascribed to ventricular fibrillation induced by neurogenic shock. Sudden immersion in icy water may act in this way extreme fright a blow over the heart insufficient to cause material damage. Two factors seem important in these instances a certain diathesis which used to be called status lymphaticus but which is probably more related to suprarenal function and a ventricle prone to fibrillation as in elderly subjects or in those with subclinical coronary artery disease. This type of death is similar to that which may be caused by a small pulmonary embolism in experiments in dogs the size of the embolism being quite insufficient to embarrass the circulation and death being preventable by atropine. The mechanism is probably a vagal reflex. It is possible that ventricular standstill may be responsible rather than ventricular fibrillation but experiments favour the latter.

Rupture of the aorta is more likely to occur from a fall especially if there is congenital hypoplasia as in many cases of coarctation. Hæmorrhage is usually into the pleural cavity or pericardium.

RUPTURE OF THE HEART

Rupture of one or more chambers of the heart following trauma is not always immediate nor does it always cause sudden death. A myocardial bruise may result in cardiac aneurysm or delayed rupture usually during the second week as described by Bright and Beck. These authors collected over 150 cases of traumatic rupture of the heart from the literature and found the incidence of the various chambers involved to be as follows:

Left ventricle	37
Right ventricle	31
Left auricle	30
Right auricle	36

More than one chamber	13
Interventricular septum	11
Interauricular septum	1

It will be appreciated that this distribution is very different from that seen with spontaneous rupture secondary to cardiac infarction when the left ventricle is nearly always responsible.

The latent interval was also studied by Warburg (1938). It occurred in 15 out of 51 cases proved at necropsy. A small tear may behave similarly to a direct penetrating wound that causes delayed death from hæmopericardium usually within a few days. A bruise may rupture at any time within six weeks (Barber 1938) or occasionally after a longer interval. Cardiac aneurysm resulting from a bruise may rupture years afterwards (Joachim and Mays 1927).

During the quiescent phase the patient may seem relatively well any discomfort being attributed to the bruise on the chest and he may continue his normal activities including sport (Priest 1939). In other cases symptoms may result from hæmopericardium or from any of the other effects to be described presently.

Diagnosis. If the patient is seen alive after cardiac rupture the signs and symptoms are those of hæmorrhage into the pericardium or pleural cavity. The combination of collapse, rapid thready pulse and a high jugular venous pressure from cardiac tamponade is very suggestive if discovered within a month of injury. There may be no evidence of external damage to the chest wall and the history of the accident may not be mentioned for it may not appear to be connected with the illness. If the possibility of previous trauma is considered the diagnosis is usually obvious.

Treatment. Immediate surgical repair is the only hope of saving life.

HÆMOPERICARDIUM

Symptoms and signs of pericarditis with or without hæmopericardium (page 355) are relatively common after indirect cardiac trauma particularly perhaps after blast injury. They provide useful evidence of cardiac damage but do not necessarily indicate its nature. Surgical interference is only warranted if there is tamponade which usually signifies cardiac rupture or serious coronary hæmorrhage. Many cases have recovered spontaneously (Smith and McKeown 1939).

MYOCARDIAL BRUISING

Crushing of the chest, direct blows over the heart and blast may all cause myocardial contusion, the clinical picture resembling that of myocardial infarction including the characteristic electrocardiographic changes or heart failure without pain (Barber 1940; Barber and Osborn 1941).

It is of considerable interest and medico-legal importance that following a direct blow in the præcordial region electrocardiographic changes may

occur which are indistinguishable from posterior myocardial infarction (Anderson 1940). Whilst it is possible that this represents remote contusion it is perhaps more likely that an anterior lesion may occlude the right coronary artery. This kind of effect will be considered more fully later.

The chief danger of myocardial contusion is delayed rupture as previously described.

Treatment consists of rest in bed for six weeks, semi starvation, a low sodium intake, mersalyl if necessary, sedatives and avoidance of digitalis.

RUPTURED AORTIC CUSP

Indirect trauma sometimes ruptures an aortic cusp. There may or may not be underlying aortic valve disease, congenital or acquired. The lesion results in the abrupt development of aortic incompetence which throws a heavy burden upon an unprepared left ventricle so that failure of that chamber is likely to ensue.

The diagnosis is suggested by the sudden onset of orthopnoea, paroxysmal cardiac dyspnoea or pulmonary oedema following a serious fall or other violent accident and is confirmed by the discovery of a loud, harsh, sometimes musical, aortic diastolic murmur often accompanied by a thrill especially if the valve was known to have been normal previously.

The prognosis may be good if the patient survives the immediate insult but death from heart failure within six weeks is a grave risk (Barber 1938, 1944). Treatment consists of six weeks' rest in bed in order to allow time for adequate compensation, and may have to be directed towards combating left ventricular failure. It must be understood that a degree of aortic incompetence which would be well tolerated and consistent with years of active life if it had developed slowly may cause death from acute heart failure when it occurs abruptly just as acute hypertension may cause left ventricular failure and pulmonary oedema whereas much higher pressures may be tolerated when developing slowly in benign hypertension.

TRAUMATIC MITRAL INCOMPETENCE

A severe fall or sudden blow over the heart or other violent accident may occasionally rupture chordae tendineae or tear one of the mitral cusps particularly if already diseased. The lesion is rare but there are many well authenticated instances (Barber and Osborn 1937). A clinical diagnosis may be made from the history if it is known that no murmur was present before the accident, if a loud harsh mitral systolic murmur is heard when the heart is first examined after the accident, if there is no evidence of previous rheumatic valve disease and if confirmatory signs of organic mitral incompetence develop, e.g. mitral systolic thrill, left ventricular enlargement and systolic expansion of the left auricle (page 282).

A number of cases have died from congestive heart failure within a few hours or weeks of the accident and others have developed mitral stenosis.

later (Barber 1938) On the other hand the accidental discovery of symptomless mitral incompetence attributable to trauma need cause little alarm such cases behaving like rheumatic mitral incompetence with a healthy myocardium

HEART BLOCK

There have been a number of instances of asphyxia in which hæmorrhage has taken place around the bundle of His with resulting heart block Several cases have been seen at necropsy by the author and a good example was observed during the 1940-1 London air raids

A woman of about 35 known to have been in previous good health was rescued in a partly asphyxiated condition from beneath a lot of debris Examination shortly afterwards revealed not only complete heart block but also gross signs of hemi Parkinsonism presumably due to hæmorrhage into the bundle of His and into the substantia nigra She declared that she had received no severe blow on her chest nor significant crush but had been partly asphyxiated by dust for about one hour

Heart block may also result from a blow over the heart or from a fall on the chest (Coffen 1930 Warburg 1938) and has been so produced experimentally in dogs (Kissane 1937) Hæmorrhage into the conducting system is presumably responsible The lesion may be transient or permanent the prognosis depending on the presence or absence of Stokes Adams fits and upon the rate of the idioventricular pace maker but on the whole it is fairly good provided there is no more serious injury and provided the heart muscle is sound

AURICULAR FIBRILLATION (OR FLUTTER)

Several cases of auricular fibrillation caused or precipitated by blows have been reported (Kahn and Kahn 1928) particularly in the elderly (Barber 1938) Bramwell (1934) records a case in which auricular fibrillation was probably initiated by a head injury and Hay and Jones (19-7) describe one due to electric shock

The mechanism whereby head injury may cause auricular fibrillation is particularly interesting though still obscure There is reason to believe that parasympathetic activity may be culpable Thus digitalis which stimulates the vagus may cause auricular fibrillation and there is a form of sinus bradycardia due to vagal influence which is associated with paroxysms of flutter or fibrillation In experiments on certain animals fibrillation may be induced by vagal stimulation Not only head injury but also meningitis Meniere's syndrome and probably other intracranial disturbances may excite this rhythm change

CARDIAC INFARCTION AND ANGINA PECTORIS

As already described myocardial contusion may give rise to clinical and electrocardiographic features similar to those of myocardial infarction and

may also result in cardiac rupture or aneurysm. There appears to be a closer relationship however between trauma and ischaemic effects. For example an anterior injury to the chest may cause a posterior left ventricular lesion clinically indistinguishable from a cardiac infarct, and classical angina pectoris may develop for the first time immediately after trauma (Campbell 1939). Moreover the subsequent course of these cases may be that of idiopathic ischaemic heart disease. It is possible that blows, crush injuries, and blast may injure the anterior coronary vessels either by causing subintimal haemorrhage in an atherosclerotic artery or more directly and thus cause acute coronary occlusion or secondary thrombosis. After such an event subsequent angina pectoris would be readily understood. Great care must be taken in diagnosing traumatic angina however for many persistent chest pains following injury represent a compensation neurosis.

Treatment consists of three to six weeks rest in bed followed by one to three months convalescence to allow time for the development of adequate collateral vascularisation. The prognosis depends upon the degree of underlying coronary disease as well as upon the amount of damage inflicted. On the whole it is not dissimilar to that in ischaemic heart disease in general.

MEDICO LEGAL ASPECTS

Employees are entitled to compensation if it can be shown that trauma has initiated or aggravated a cardiovascular disability. Even a case of syphilitic aneurysm that ruptures during the course of work receives compensation. Patients with established heart disease may deteriorate after an accident and this aggravation is equally compensated. The benefit of doubt is always given to the patient and in a court of law or a tribunal it is difficult to convince a judge or president that trauma has not adversely affected the cardiovascular system. Yet a firm stand must be taken over the development of cardiac neurosis. Left inframammary pain is especially liable to become persistent and intractable if linked to the idea of compensation and the physician must be prepared to make a categorical statement to the effect that this is not organic and is not due to the accident, that its origin lies in the mind and in the emotions and its growth runs parallel with the conscious or subconscious desire for gain.

REFERENCES

SPONTANEOUS LESIONS

Abbott M L (1919) Clinical and developmental study of a case of ruptured aneurysm of the right anterior aortic sinus of Valsalva. Contributions to medical and biological research. New York.

Baer S and Goldburgh H L (1948) The varied clinical syndromes produced by dissecting aneurysm. *Amer Heart J* 35 198.

David P McPeak E M Vivas Salas E and White P D (1947) Dissecting aneurysm of the aorta review of 17 autopsied cases of acute dissecting aneurysm of the aorta encountered at the Massachusetts Gen Hosp from 1937-46 inc eight of which were correctly diagnosed ante mortem *Ann intern Med* 27 405

Frdheim J (19 9) Medionecrosis aortae idiopathica *Irish Arch f path Anat* 273 454

Flaxman A (1942) Dissecting aneurysm of aorta *Amer Heart J* 24 654

Gager L (19 8) Dissecting aneurysm of aorta complicating hypertension *Ibid* 3 489

Galbraith A J Gardner F and Hardwick S (1939) Huge dissecting aneurysm *Lancet* ii 1019

Gouley B A and Anderson E (1940) Chronic dissecting aneurysm simulating syphilitic cardiovascular disease notes on associated aortic murmurs *Ann intern Med* 14 978

Hamburger M and Ferris E B (1938) Dissecting aneurysm *Amer Heart J* 16 1

Lænnec R T H (1826) *Traite de l'auscultation mediate* 2nd Edit Vol II 696 3rd Edit Vol III 295

Lawrence J H (1935) Clinical symptoms and signs of dissecting aneurysm of aorta with report of case diagnosed during life *Internat Clin* 2 122

McGeachy T E and Paullin J E (1937) Dissecting aneurysm of aorta *J Amer med Ass* 108 1690

Moritz A R (1932) Medionecrosis aortae idiopathica cystica *Amer J Path* 8 717

Mote C D and Carr J L (1942) Dissecting aneurysm of the aorta *Amer Heart J* 24 65

Peacock T B (1863) Report on cases of dissecting aneurysms *Trans path Soc London* 14 87

Peery T M (194) Incomplete rupture of the aorta *Arch intern Med* 70 689

Rottino A (1939) Medial degeneration of aorta as seen in 12 cases of dissecting aneurysm *Arch Path* 28 1

Schnikter M A and Bayer C A (1944) Dissecting aneurysm of the aorta in young individuals *Ann intern Med* 20 486

Shennan T (1934) Dissecting aneurysms MRC report London

Taussig H (1947) Congenital malformation of the heart Commonwealth Fund New York

Tyson M D (1931) Dissecting aneurysms *Amer J Path* 7 581

Wainwright C W (1944) Dissecting aneurysm producing coronary occlusion by dissection of coronary artery *Bull Johns Hopk Hosp* 75 81

Weisman A D and Adams R D (1944) Neurological complications of dissecting aneurysm *Brain* 67 69

Weiss S (1935) Clinical course of spontaneous dissecting aneurysm of aorta *M Clin N Amer* 18 1117

Wood F C Pendergrass E P and Ostrum H W (193) Dissecting aneurysm of aorta with special reference to its ro ntgenographic features *Amer J Roentgenol* 28 437

EFFECTS OF DIRECT OR INDIRECT INJURY

Ander on R G (1940) Non penetrating injuries of the heart *Brit med J* ii 307

Barber H (1938) Trauma of the heart *Ibid* i 433 — (1940) Contusion of the myocardium *Ibid* ii 520 — (1944) The effects of trauma direct or

- indirect on the heart *Quart J Med* 13 137 — Osborn G R (1937)
 Case of mitral stenosis result of trauma *Guy's Hosp Rep* 87 510 — —
 (1941) A fatal case of myocardial contusion *Brit Heart J* 3 127
 Bramwell C (1934) Can a head injury cause auricular fibrillation? *Lancet* 1
 8
 Bright E F and Beck C S (1935) Non penetrating wounds of the heart
 a clinical and experimental study *Amer Heart J* 10 293
 Campbell M (1939) Angina pectoris following a crushing accident *Brit
 Heart J* 1 177
 Coffen T H (1930) Complete heart block of 7 years duration in child
 resulting from injury *Amer Heart J* 5 667
 Hay J and Jones H W (1927) Trauma as a cause of auricular fibrillation
Brit med J 1 559
 Joachim H and Mays A T (1927) A case of cardiac aneurysm probably
 of traumatic origin *Amer Heart J* 2 682
 Kahn M H and Kahn S (1929) Cardiovascular lesions following injury to
 the chest *Ann intern Med* 2 1013
 King F S J (1941) Surgery of the heart London
 Kissane R W (1937) Contusion of the heart Columbus
 Nicholson W F (1945) War wounds of the heart Conf Army Phys Rome
 Priest R (1939) Notes on three interesting cases I Trauma of the heart
J Roy Army Med C 73 125
 Smith L B and McKeown J H (1939) Contusion of the heart *Amer
 Heart J* 17 561
 Turner C C (1941) A bullet in the heart for twenty three years *Surg* 9 83-
 Warburg E (1938) Traumatic heart lesions London
 Wood P H (1937) Electrocardiographic changes of a T₂ pattern in pericardial
 lesions and in stab wound of the heart *Lancet* ii 796 — (1945) War wounds
 of the heart Conf Army Phys Rome

CARDIOVASCULAR DISTURBANCES ASSOCIATED WITH PSYCHIATRIC STATES

THE cardiovascular system may be profoundly influenced by psychological or psychiatric states through the medium of the autonomic nervous system. The stimulus is emotional and appears to act on the central vegetative nuclei in the region of the hypothalamus. We are all familiar with the uncomfortable thudding of our hearts during moments of fear and most of us have witnessed a fainting attack provoked by the sight of something which is at once queer and frightening. The physiological basis for such phenomena is relatively simple: sympathetic or adrenergic activity may cause palpitations by accelerating the pulse, elevating the blood pressure and strengthening the heart beat; parasympathetic or cholinergic activity may induce syncope by retarding the pulse, lowering the blood pressure and weakening the heart beat.

Cardiovascular upsets of this kind, sufficient to bring the patient to seek medical advice, almost invariably indicate psychiatric disorder, for the effects of emotion within the limits of common physiological experience are too transient and too familiar to disturb a normal individual. Moreover, in psychiatric states such symptoms may be persistent or may be provoked too readily. The syndrome so produced has been called soldier's heart, irritable heart, disordered action of the heart (D.A.H.), cardiac neurosis, effort syndrome, autonomic imbalance, neurocirculatory asthenia, etc. Such terms should be discarded in favour of the correct psychiatric diagnosis, but the words effort intolerance may be added with advantage, preferably in brackets, when clinically important. Historically one may speak of Da Costa's syndrome to cover all previous nomenclature (Wood, 1941).

The syndrome is characterised by a group of symptoms which unduly limit the subject's capacity for effort or which upset his peace of mind at rest by a number of signs which depend upon disturbance of the autonomic nervous system and by an underlying psychiatric disorder. The cardinal symptoms are breathlessness (93 per cent), palpitations (89 per cent), fatigue (88 per cent), left inframammary pain (78 per cent) and dizziness (78 per cent) or syncope (35 per cent). The cardinal signs are those of functional disturbance of the respiratory, vasomotor, sudomotor and muscular systems. The psychiatric disorder is commonly an anxiety state, but may be almost anything with high emotional content, including the psychoses.

It should be understood that there is no essential difference between effort syndrome and cardiac neurosis: they are merely clothed differ-

ently the former in battle dress the latter in artificial silk. In civil life the condition accounts for 10 to 15 per cent of all cases referred to cardiovascular clinics: it is common in children, and occurs more often in women than in men the ratio being 3:2. It has a preference for the emotional races especially the Jews and the Italians. In the first world war there were some 60 000 effort syndrome casualties in the British forces: in the second a more enlightened view was taken the majority of these cases receiving appropriate psychiatric labels and management.

CLINICAL FEATURES

The cardinal symptoms and signs have already been mentioned they will now be discussed in more detail.

Breathlessness These patients experience a true sensation of breathlessness in circumstances that would not affect a normal person. It is not only a question of breathlessness on effort but patients will say they are unable to obtain a satisfying breath or that they feel a sense of suffocation and this is confirmed objectively by frequent deep sighs. Sometimes they complain of attacks of nocturnal dyspnoea which may be confused with bronchial asthma or with paroxysmal cardiac dyspnoea: careful questioning however should reveal their psychosomatic nature especially by probing the precipitating anxiety dream and by unmasking the associated panic state. Further evidence of functional respiratory disorder may be obtained by noting hurried irregular and shallow breathing. A simple and illuminating test is forced hyperventilation. The patient is asked to breathe deeply and rapidly for one minute. A normal individual experiences dizziness and sometimes slight tingling of the fingers and toes. When told to stop he passes into a state of apnoea lasting about 20 seconds. The psychoneurotic especially the hysteric dramatises his subjective sensations and when told to desist usually continues forced breathing explaining later that he felt breathless. Since dizziness and tingling of the extremities are due to vasoconstriction induced by carbon dioxide washout it is clear that such psychoneurotics experience breathlessness when the carbon dioxide content of the arterial blood is so low as to cause apnoea in controls. The respiratory stimulus must therefore come from higher centres. The maximum breath-holding time is another useful test. Normal subjects have no difficulty in holding the breath for at least 30 seconds but patients with Da Costa's syndrome usually give up very quickly 30 per cent of them in less than 10 seconds moreover in contrast to controls they show little distress when they reach the breaking point.

Palpitations Cardiac overaction resulting from emotional stimulation plays an important role in the induction of cardiac neurosis. It is a common psychiatric event for some intangible fear to become linked to something more easily understood and remote from the real difficulty. For example a psychoneurotic with a morbid fear of heights may develop palpitations

when ordered to climb a ladder. If the idea that palpitations may denote some disorder of the heart occurs to him, he at once embraces the possibility and proceeds to advance the theory in all seriousness, for it disguises his true fear which might be thought shameful and protects him from the danger. Although a successful defence mechanism in these two respects the manoeuvre is baneful because it provokes a new fear, that of heart disease and sudden death, this new fear aggravates the palpitations and so closes a vicious circle.

The palpitations of anxiety states are associated with sinus tachycardia, elevation of the blood pressure, increase in cardiac output and probably with strengthening of the heart beat. These features are due essentially to emotional stimulation of a normal adrenergic system.

Fatigue. Patients often complain that they do not feel refreshed when they wake in the morning, that their sleep has been of no benefit to them. They also feel tired and listless during the day and are unduly fatigued by effort. The symptom is usually attributed to anxiety dreams and to emotional conflicts.

Left inframmary pain. Psychosomatic pain is usually situated in the left inframmary region, but may be higher, lower, more central or more lateral; it may radiate down the left arm. It is commonly described as aching or as sharp and stabbing in quality, but occasionally it is constricting or cramp-like. Although pain may occur during effort, it is more frequent afterwards; it is also common at night and may prevent the patient sleeping on the left side. Sometimes it is capricious and bears no relationship to any known factor. Sharp twinges are momentary and acute stitch-like pain may last several minutes, but the classical ache usually continues for hours. It thus usually differs from angina pectoris in its eccentric site, in its quality, in its relationship to effort and in its duration, i.e. in every important respect. Occasionally, however, as may be inferred from the description given above, psychosomatic pain may be situated near the left border of the sternum, referred to the left arm, constricting in quality and measured in minutes. In such cases it may well be misinterpreted. There is usually some odd remark, however, or something in the patient's manner, which should warn the physician and encourage him to launch a critical cross-examination. The precise history of angina pectoris will not be shaken by this, but that of an anxiety state alters and becomes more complicated and confused when elaborated.

Left inframmary pain is important because it seems to convince the patient that his heart is diseased and it is not unnatural that he should think thus of a pain arising so close to it. In the psychoneurotic this creates a morbid fear of death and catastrophe and so closes another vicious circle.

The exact mechanism of the pain is obscure. It is immediately abolished by the intramuscular injection of 2 ml. of novocaine at the site of maximum intensity or tenderness. Cutaneous or subcutaneous anaesthesia has no

effect This indicates that it is not referred but arises locally in muscle or fascia and suggests that it is related to fibrositis and low back pain It may be initiated by fatigue or strain of respiratory muscles in cases with respiratory neurosis by strain of certain muscular attachments involved in such actions as cranking an engine or lifting a heavy weight by incessant minimum trauma from the light hammer blows of an overacting heart or by faulty posture It is exaggerated and perpetuated by the belief that it arises in the heart

Dizziness Dizziness means momentary faintness, transient unsteadiness, light headedness or a far away feeling It does not refer to spinning as in vertigo It may occur on sudden movement of the head on standing, up abruptly or during effort It is readily reproduced by hyperventilation when it is attributed to cerebral vasoconstriction Orthostatic dizziness is related to orthostatic hypotension and is due to inadequate circulatory adjustments on assuming the erect posture It is probable that other forms of dizziness are also due to diminished cerebral blood flow induced by autonomic disturbance Transient loss of consciousness due to temporary failure of the cerebral circulation occurs at one time or another in 20 to 30 per cent of these cases

Sweating Sweating is a helpful diagnostic feature because in the majority of instances it is confined to the axillæ to the palms of the hands and to the sole of the feet These are emotional sweat areas Thermal sweating and that induced by cholinergic drugs have a different distribution being much more widespread Sweating associated with effort may begin emotionally but is soon thermal Thyrotoxic sweating is also thermal The hands are the best single guide if sweating is confined to the palms the stimulus is emotional if the backs of the hands are also involved other causes should be considered Undue sweating is mentioned or admitted by 80 per cent of these cases and is seen objectively in about two thirds

Headache Headache is a common complaint (72 per cent) and is either vague or throbbing In assessing the reality of the physical basis of the throbbing type it is helpful to ask the patient to count the throb aloud or better to tap out the rhythm digitally while the observer checks this against the pulse rate in true vascular headache they must coincide in hysteria they do not Unilateral carotid compression is also useful for it abolishes vascular headache on the same side but it either aggravates or has no effect upon hysterical pain Throbbing vascular headache may be induced by the intravenous injection of 1 mg of histamine or by trinitrin or amyl nitrite in some cases It is closely associated with exaggerated pulsation of the cerebral arteries (Pickering 1939) It is seen clinically not only in the anxiety states but also in fevers and in acute alcoholism It occurs spontaneously in migraine Improvement depends upon better autonomic regulation which in turn depends upon successful treatment of the underlying anxiety state



Fig. 21.01—Classical face, build and posture of a case of Dress Addict's Syndrome. Painted by Ian Tillaud (life-size portrait in the museum of the Post Graduate Medical School of London)

PHYSICAL SIGNS

Signs of autonomic disturbance serve to check the validity of psychosomatic symptoms. Most have already been mentioned but they will be recapitulated and grouped here for convenience.

General

- Tense dejected or diffident manner
- Dull weak or listless facies
- Soft quiet timid voice

Cardiovascular

- Tachycardia (30 per cent)
- Overaction of the heart (44 per cent)
- Blood pressure in the region of 150/90 mm Hg (27 per cent above)
- Deceleration time over 2 minutes in effort tolerance test (33 per cent)
- Acrocyanosis (44 per cent)
- Flushes (36 per cent)

Respiratory

- Frequent deep sighs (32 per cent)
- Rapid irregular or shallow breathing occasionally hyperventilation (21 per cent)
- Inability to hold the breath for 30 seconds (76 per cent)
- Dyspnoea instead of apnoea after forced breathing

Sudomotor

- Visible sweat on the palms of the hands (67 per cent)
- Sweat trickling from the axillae (35 per cent)

Skeletal and Muscular

- Tremor of fingers usually coarse irregular and inconstant (26 per cent)
- Shakiness of voice and limbs
- Asthenic posture or poor physical development (41 per cent)
- Tenderness in area of left inframmary pain

A life sized portrait of one of these patients (fig. 21.01) hangs in the library of the Postgraduate Medical School of London and surpasses any description. The effort tolerance test consists of stepping on and off a chair ten times and counting the pulse rate before immediately after and subsequently at minute intervals until the resting speed is regained. The deceleration time is abnormal (over 2 minutes) in 33 per cent of these patients.

Physical signs of autonomic disturbance are helpful in distinguishing the malingering and in assessing the severity of the case. About 90 per cent of normal young adults do not show more than one of these signs and 50 per cent show none.

PSYCHIATRIC ASPECTS

Although the syndrome described may occur in any psychiatric state with high emotional tone it is usually associated with an anxiety state. In many there are hysterical features and a large number show reactive depression.

The family history is tainted with psychoneurosis in 50 to 60 per cent compared with 5 to 10 per cent in controls with or without organic heart disease. About 66 per cent describe neurotic traits in childhood: morbid fears especially of the dark, of heights, of water or of animals are frequent; bed wetting, stammering, tics, nightmares, sleep walking and undue delicacy of health are common. They are timid children, far too dependent upon maternal protection. At school, kindly doctors and soft mothers protect them from the hazards of football, swimming and the gymnasium.

It is probable that predisposition to psychoneurosis is mainly hereditary but early environmental factors such as domestic strife, insecurity, suppression and maternal coddling play their part.

There are many factors which may operate to bring about the adult syndrome and in any particular case one should never be satisfied with the discovery of only one or two. It is fruitful to search for evidence of predisposition for a state of mind recently prepared for the development of psychoneurosis by external or by endogenous factors, for precipitating agents for the growth of vicious circles and for motives for gain that aggravate and perpetuate the syndrome. Proper assessment, management and prognosis are impossible if any vital link is overlooked.

Hereditary and environmental predisposition have already been discussed. The mind is especially prepared for the development of psychoneurosis when in a state of confusion and unreality. Head injuries effect this; certain acute fevers are often responsible, especially rheumatic fever, influenza, meningitis and diphtheria; long hours of work in unpleasant and unhappy surroundings may be to blame.

Precipitating factors are often multiple. It is as if one or two could be coped with but when several occur one on top of the other, mental equilibrium disintegrates. They are usually closely linked with fear in some form or another. The most obvious example is active service; hence the high incidence of the disorder in war. Fear of football and fear of swimming are common in childhood and may precipitate anxiety at school. The fear of being unsuccessful, of not being able to shoulder responsibility is a common cause of breakdown in civil life. Insecurity or fear of the future is also common. The adoption of a line of action contrary to established social custom may cause an anxiety state due to fear of discovery and public criticism. Difficult personal relationships, especially between husband and wife, are often responsible. Sex difficulties are important but should not be over emphasised. Financial worry, unemployment and fear of disease play their part. To a timid sensitive character the fear of

being found out of being thought a coward of being proved inadequate of seeming a fool—and so of losing cast is a very real and powerful emotional stimulus

The development of vicious circular patterns is interesting. In this particular syndrome most vicious circles have a common basis and revolve round the fear of heart disease and sudden death. The combination of breathlessness, dizziness or syncope, fatigue and especially palpitations and left inframammary pain provides convincing evidence of heart disease to the lay mind. All these symptoms, which are psychosomatic in mechanism, may be produced by simple anxiety and may disappear rapidly as soon as the anxiety is resolved. But if the patient takes the fatal step and believes that they are due to heart disease, a vicious circle is at once established. For a new and greater fear develops, that of sudden death at any moment. This constant anxiety, operating consciously or subconsciously every second of the day and night, increases the severity of the psychosomatic symptoms. Under these circumstances the syndrome is maintained long after resolution of the original anxiety. Superimposed upon this pattern or independent of it, there develop various and often complicated conditioned reflexes, until finally distressing autonomic reactions are so ingrained and so divorced from conscious thought as to be practically ineradicable. Correct medical interpretation of early psychosomatic symptoms is of the utmost importance in the prevention of these pernicious grooves. The doctor who misinterprets a boy's fear of water and accepts the pallor and palpitations as signs of heart disease, who mistakes left inframammary pain for angina pectoris, who finding an innocent systolic murmur diagnoses valvular heart disease, who regards syncope or dizziness as a sign of cardiac weakness, is guilty not only of stupidity and ignorance, but is also responsible for turning his patient into a chronic and incurable psychoneurotic. Even so, it may be comforting to know that medical blunders of this kind will influence only 10 per cent of apparently normal individuals, the great majority adversely affected showing evidence of predisposition.

Finally, there is the motive for gain. This is seen in compensation neurosis and in war it is obvious at every medical board. The inadequate personality of so many of these patients capitalises the symptoms. What timid man, indifferent to higher ideals, will face the dangers of battle when the very symptoms of his fear offer him protection?

DIFFERENTIAL DIAGNOSIS

The characteristic symptoms and signs associated with psychiatric disorder usually make the diagnosis easy. The physical features have been stressed because the psychiatric state may not be obvious until the mind has been deeply probed. This is well shown by comparing the conclusions drawn at the special investigation centres for effort syndrome during the

first two world wars at Hampstead in world war I where little attention was paid to psychiatry not more than 10 per cent were considered psychoneurotic at Mill Hill in world war II a psychiatric basis was proved in 94 per cent The diagnosis should be positive not dependent upon a process of exclusion it may stand even when organic disease is also found especially mild rheumatic heart disease benign hypertension and chronic bronchitis

Thyrotoxicosis may present difficulty to the inexperienced The common mistake is to diagnose an anxiety state as thyrotoxicosis rarely the reverse The difference is fully considered on page 481 and 485 Particular attention should be paid to the attitude and behaviour of the patient to the expression of the eyes to the colour and temperature of the hands to the distribution of sweating to the diastolic blood pressure and to the appetite

In children active rheumatic carditis may cause confusion vague muscle pains being mistaken for joint pains and tics for chorea

Attacks of violent palpitations in anxiety states are sometimes confused with paroxysmal tachycardia Accurate history taking and observation of an induced attack should prevent error The special points of difference are given on page 133

The distinction between left inframammary pain and angina pectoris has already been considered but real difficulty may arise In both the diagnosis depends largely upon the history and cannot be proved or disproved by the demonstration of psychoneurosis on the one hand or of organic heart disease on the other The matter is further complicated by the adverse effect of anxiety upon ischæmic heart disease for it may be so important a factor that its satisfactory resolution may temporarily relieve angina pectoris Occasionally the diagnosis remains doubtful until determined by the future course

The physician should be on his guard against pulmonary tuberculosis chronic undulant fever juvenile spondylitis spontaneous hypoglycæmia subacute bacterial endocarditis deficiency of the vitamin B group and certain endocrine disorders—especially the menopause Anæmia should be more obvious When the symptoms first arise during convalescence simple reassurance should be given and the final diagnosis deferred until it is clear that rapid recovery has or has not taken place

TREATMENT

Treatment is never easy and is the more difficult the longer it is delayed Failure is certain if any essential factor in the development of the syndrome is overlooked so that a great deal of time must be spent on these patients Simple reassurance and some superficial explanation are quite inadequate

First the patient must feel that at last he has met a doctor who thoroughly understands his case secondly a complete physical examination supported

by fluoroscopy and an electrocardiogram is necessary so that he will respect unconditional reassurance. Adequate explanation must follow, and will vary according to the chief symptoms. The object is to convince the patient that the symptoms are emotionally produced. One may point out how sudden fear causes palpitations, sweating, alteration of breathing and sometime a fainting attack. He will agree with this but may object that he feels no such fear. One should then explain that great fear acting for a few seconds may be more than equalled by a tiny remote fear acting over weeks, months or years, a state called anxiety. This step is difficult, but the point must be carried. Correct interpretation of anxiety dreams is of value in demonstrating the power of subconscious emotion. Insight and conviction may come suddenly if psychosomatic disturbance on some particular occasion or under certain specific circumstances can be explained in the light of emotional experience.

For example, a patient at Mill Hill gave a history of a morbid fear of fireworks in his boyhood, conditioned by London air raids in his infancy. Otherwise he was fit and strong. He was called up in September 1939, was sent to France and remained well until told one day to unload an ammunition lorry. On handling the shells he became curiously panic stricken, developed gross psychosomatic symptoms and misinterpreted them, thinking they meant heart disease. A vicious circle was initiated, he reported sick, and finally arrived at a base hospital with an established effort syndrome. When the link between his fear of handling fireworks and his handling shells for the first time was pointed out, he was suddenly convinced of the truth of the explanation given for his symptoms and made a rapid and complete recovery. But his fear of fireworks, shells and all other explosives was unabated. Treatment had only been directed towards the removal of effort intolerance, by abolishing the misinterpretation and vicious circle that initiated and maintained it.

As a rule, however, it is not enough to reassure and give an adequate explanation, for by the time the patient consults a physician the syndrome is usually highly complex and conditioned reflexes are well ingrained. To cut across such reflexes and vicious circles, one may encourage the patient to come to better terms with his symptoms. He fears them because he thinks they are injurious and may result in sudden death. He must be told they are harmless, that they can never be more than a nuisance, that he is already familiar with the worst they can do. Once he appreciates the fact that if he no longer fears his symptoms he will cease to aggravate them, the point is scored.

If there is an hysterical motive for gain, it must be mentioned and then ruthlessly underlined. It is remarkable what little insight these patients have and disconcerting how little shame.

The methods so far outlined do not touch the underlying psychoneurosis and the real treatment has yet to begin. The patient may be referred to a psychiatrist or if the causative factors seem clear the physician may prefer to deal with them himself. There are always three things to consider: the

difficulties in which the patient is floundering his reaction which is based on his character and intelligence and his attitude towards his reaction. The difficulties should be taken first sorted out and resolved as far as possible. The help of social welfare workers may be enlisted in this respect. The patient's reaction should be analysed and some psychiatric skill and knowledge are required to do this. It is often possible to show that his reaction is based on false values, ideas or beliefs. Or one may simply explain just why he so reacts in order to give him insight. It is impossible to outline precisely just what is required for every case is different and needs individual treatment. If the problem has no satisfactory solution and if the patient's reaction cannot be altered favourably then at least he may learn to get on better terms with both. Difficulties must be faced and not hidden away in the dark recesses of the mind. Highly personal matters should be fully discussed in a matter of fact way until they cease to seem so dreadful if a man is standing on a false pedestal he must learn humility and honesty and tread upon the good earth.

Finally the background must be assessed. With strong hereditary taints and bad early environment the outlook is poor and the aim should be to fit the patient into circumstances which will cause the least embarrassment. This is a confession of failure. At the other extreme if the stock is good and if there is no evidence of predisposition and if this is confirmed by the severity of the stress of anxiety causing the breakdown every effort should be made to cure the patient. In other words one should deal with the environment when the prognosis is bad and with the patient when it is good.

REFERENCES

- Pickering G W (1939) Experimental observations on headache *Brit med J* 1: 907
 Wood P H (1941) Differential diagnosis of Da Costa's syndrome *Proc Roy Soc Med* 34: 543 — (1941) El síndrome de Da Costa *Archiv Latino Americanos Cardiol Hematol* 11: 41 — (1941) Da Costa's syndrome *Brit med J* 1: 767 805 845
 (A full bibliography is contained in these three articles)

INDEX

- a wave (venous) 170
 - in auricular fibrillation 13
 - in heart block 122
 - in heart failure 170
 - normal 19
- a wave (venous) giant
 - in pulmonary hypertension 462
 - in pulmonary stenosis 229
 - in tricuspid incompetence 304
 - in tricuspid stenosis 229
- A C interference 68
- Abdominal compression and venous pressure 169 171
 - in paroxysmal hypertension 421
- Accelerator nerves 109
- Accession wave 65
- Acetyl B methyl choline (mécholin) 138 200 316
- Acetyl choline 138 199
- Acidosis 200 464
- Acrocyanosis 540
- A C T H 275
- Acupuncture 189
- Acute (see item required)
- Adams Stokes syndrome (see *Stokes Adams*) 122
- Adenoma fetal of thyroid 486
- Adenomatous goitre 479 486
- Adherent pericardium 351
 - rheumatic 308
- Adhesions pleurop pericardial 165
- Adrenal
 - action of 111 205 326 462
 - in adrenalm dulatorytumour 417 421 422
 - in allergic reactions 338
 - in angina pectoris 326
 - in beri beri 516
 - in bronchial asthma 473
 - in cardiac asthma 190
 - in circulatory failure 314
 - in heart block 124
 - insensitivity in myxoedema 496
 - in myocardial infarction 190 407
 - overdosage 326
 - in paroxysmal hypertension 421
 - in Stokes Adams syndrome 124
 - and ventricular fibrillation 150
- Adrenal cortical tumour 436
- Adrenal medullary tumour 421-2
- Air complementary 17
 - serv 17
 - residual (see *Emphysema*) 17
- Air embolism 457
 - in cardiac catheterisation 16
- Albuminuria 21
 - in heart failure 21 22 174
 - in hypertension 433
 - in nephritis 433
- Alcohol 326
 - in angina pectoris 371 381
 - in beri beri 514
- Alkaline phosphatase 517
- Alkalosis in hyperventilation 200 536
 - vaso constriction in 536
- Allergic myocarditis 314 315 318
- Allergic shock 311
- Alphatocopherol (vitamin E) 384
- Alternation
 - of the pulse 166
 - of the QRS complex 167
- Altitude anoxæmia and 201 3 4
- Aminophylline in angina pectoris 38
 - in asthma 473
 - in cardiac infarction 407
 - in Cheyne Stokes breathing 190
 - in left ventricular failure 190
 - in pulmonary embolism 455
- Aminothiazol 492
- Amni visnaga 382
- Ammonium chloride 184
- Amphibrachic metre 164
- Amyl acetate 13
- Amyl nitrite
 - action of 382
 - in angina pectoris 381
 - in diagnosis of murmurs 8 224 -88
- Anærotic pulse 3 300
- Anæmia 2- 502
 - and angina pectoris 374 503
 - in bacterial endocarditis 331 506
 - cardiac output 155 156 503
 - circulation 503
 - clinical features 503
 - diagnosis 506
 - electrocardiogram 504
 - fatty degeneration of heart 502
 - heart in 502
 - heart failure in 155 503
 - murmurs 503
 - in myxoedema 496
 - oedema 503
 - pulmonary oedema 503
 - in rheumatic fever 259
 - treatment 507
 - nous pressure 155 503
 - X ray 506
- Anæsthetic meter 165
- Anæsthesia (see *Edema*) 72

- Anatomy of
 aortic valve 61
 conducting system 108
 coronary arteries 387
 heart chambers 25 26
 left atrial appendix 216
 pulmonary arteries 14 32
 pulmonary valve 242
 pulmonary veins 33 235
 surface markings 9
- Aneurin (thiamine) in beri beri 515
 in thyrotoxicosis 491
- Aneurysm (and see *Syphilitic aneurysm*)
 arteriovenous 511
 berry 211
 cardiac (see *Cardiac aneurysm*) 44 397
 404 529
 circoid (congenital) 511
 mycotic 333
 of pulmonary artery 475
 syphilitic 359-64
- Aneurysmal dilatation of
 left auricle 284
 pulmonary artery 212 218 475
- Angina innocens 377
- Angina pectoris (and see *Ischaemic heart disease*) 374 86
 in adrenalin overdose 326 408
 in anaemia 503
 in aortic incompetence 366
 in aortic stenosis 299 366
 in auricular fibrillation 147
 in auricular flutter 142
 blood pressure in 375
 clinical features 375 378
 contributory causes 374
 course 380
 decubitus 322 376
 diagnosis 376 377
 diet 381
 effort test 378
 electrocardiogram 378
 etiology 370 374
 in hypertension 375 429
 hypoxic test 378
 in isolated myocarditis 318
 in mitral stenosis 294
 in myocardema 497
 pain 375-6
 in paroxysmal tachycardia 132
 pathology 372
 physiology 374
 prognosis 380
 in pulmonary stenosis 229
 at rest 384 408
 sex incidence 370
 sudden death 150
 surgical treatment 385
 in syphilitic aortitis 366
 thiouracil for 385
- Angina pectoris (contd.)
 in thyrotoxicosis 495
 traumatic 532
 treatment 322 380 5
 in women 370
- Angiocardiography 31
 in aneurysm of aorta 360 362 363
 in arterio-venous aneurysm of lung 513
 in coarctation of aorta 209
 in Eisenmenger's complex 245
 in Fallot's tetralogy 238 242
 in mediastinal tumour 362
 in mitral stenosis 289 290
 in patent ductus arteriosus 224
 in pulmonary atresia 250
 in pulmonary stenosis 234
 in superior vena cava obstruction 360
 in syphilitic aortitis 364
 in transposition 250
 in tricuspid atresia 250
- Angioneurotic oedema 173 338
- Angiotonin 419
- Anions 65
- Anomalies - congenital 203-251
- Anomalous left coronary artery 205
- Anoxaemia
 cardiac output 472
 electrocardiogram 101
- Anoxic
 pulmonary heart disease 464-74
 syncope 201
 tachycardia 111
 test for angina 378
- Antero-posterior diameter of the heart 34
- Antibiotics 336 9
- Anti-coagulants in
 acute coronary insufficiency 384
 bacterial endocarditis 336
 cardiac infarction 406
 thrombo-embolic states 454
- Anti-clockwise rotation 73 83 85
 in hypertension 426
- Anti-histamine drugs in
 asthma 473
 penicillin urticaria 338
 streptomycin vertigo 339
- Anti-spasmodics 455 473
- Anti-streptococcal haemolysin titre 257
 262
- Anxiety state (see *Cardiac neurosis*) 535-45
- Aorta (and see *Aortic*)
 abnormalities (A. rarsa) 38
 atherosclerosis 373
 calcification 38 39 358 365 374
 coarctation 39 206-12
 dextro position 236 244
 dilatation 38
 dissection 521
 hypoplasia 42
 kinking 474

- Aorta (contd.)**
 medial necrosis 521
 prominence 38 298 429
 riding 236 244
 rupture 528
 tortuosity 39
 unfolding 39 298 429
 X ray 38-42
 width 35
- Aortic aneurysm (see *aneurysm*)** 38 359-63
- Aortic arch** development of 206
 double 207
 right sided 42 236 242
- Aortic area** 9
- Aortic atresia** 207
- Aortic cusps** rupture of 530
- Aortic diastolic murmur (and see *Aortic incompetence*)** 295
 functional 424 503
 in rheumatic carditis 270
- Aortic diastolic thrill** in
 bacterial endocarditis 331
 perforated cusp 331
 ruptured cusp 530
 syphilis 364
- Aortic incompetence**
 in anaemia 503
 angina in 366
 atherosclerotic 303
 bacterial 330
 clinical features 295-8 364
 in coarctation of the aorta 209
 congenital 212
 diagnosis 298
 in dissecting aneurysm 523
 in Eisenmenger's complex 244
 electrocardiogram 298
 functional 424 503
 hypertensive 424
 pathology 294 364
 physiology 294 295
 prognosis 299
 pulse 296 297
 rheumatic 270 294-9
 syncope in 194 196
 syphilitic 364-366
 traumatic 530
 X ray 298
- Aortic isthmus** 207
- Aortic knob** 26
- Aortic knuckle** 26
 double 209
- Aortic notch** 20
- Aortic pressure curve** 295
- Aortic regurgitation (see *Aortic incompetence*)** 294 9
- Aortic ring** 364 424
- Aortic sinus perforation** of 229 523 5
- Aortic stenosis** 299-303
 angina pectoris in 299-300
 atherosclerotic 303
 blood pressure 301
 calcific 61 302 3
 clinical features 299-301
 congenital 212
 electrocardiogram 301
 etiology 302-3
 pathology 299
 physiology 299
 prognosis 303
 pulse 300
 rheumatic 299
 syncope in 194 299
 X ray 301
- Aortic systolic murmur**
 in aortic incompetence 364
 in aortic stenosis 301
 functional 281-2 487 503
- Aortic systolic thrill** 301
- Aortic valve**
 anatomical site 61
 bicuspid 209
 calcification 61 301
 disease (see *Aortic incompetence and stenosis*)
 vascularity 258
- Aortic valvulitis** 240
- Aortitis (syphilitic)** 358-68
- Aortography** 209
- Aorto pulmonary septum** 206
 defect of 204 523
- Apex beat (and see *Cardiac impulse*)**
 definition 6
 displacement 6
 fixation 351
- Apical systolic murmur** 273 280-2
- Apnoea**
 in Cheyne Stokes breathing 167
 post hyperpnoea 536
- Arachnodactyly** 216
- Arborisation block** 90
- Arm to lung time** 13 163
- Arm to tongue time** 12 31 161
- Arrhythmia sinus** 109
- Arsenic** 367
- Arteries** palpation of 3
 structural changes of 419
- Arterial oxygen saturation** on 17
 in cyanotic states 177
 in Eisenmenger's complex 245
 in Fallot's tetralogy 242
 in heart failure 190
 in massive pulmonary embolism 455
 in mitral stenosis 285
 in pulmonary heart disease 462 47-
 in reversed interatrial shunt 234
 rise of in oxygen tent 242
- Arterial puncture** 17
- Arterial pulse waves** 20

- Arteriogram 20
- Arterio sclerosis (see *Atherosclerosis*) 372-374 419
- Arterio venous aneurysm 511-14
- acquired 513
 - cerebral 511
 - congenital 511
 - peripheral 511
 - pulmonary 513
 - traumatic 513
 - treatment 514
- Arterio venous shunt (in) 513-14
- arterio venous aneurysm 514
- atrial septal defect 216 218
- beri beri 516
- congenital heart disease 204
- hepatic disease 517
- Paget's disease 517
- patent ductus 223 37
- pregnancy 507
- thyrotoxicosis 486 493
- ventricular septal defect 221
- Arthus phenomenon 258
- Aschoff node 258
- Ascites
- in heart failure 172 174
 - in Pick's disease 349
 - in tricuspid disease 306
 - venous pressure in 171
- Asphyxia electrocardiogram in 102
- Asphyxial hæmorrhage 531
- Asthenic posture 540
- Asthma
- bronchial 464
 - cardiac 160
 - treatment 473
- Atheroma 372 4
- Atherosclerosis 372-4 419 433
- Athlete's heart 35 110
- Atresia
- aortic 207
 - pulmonary 237 250
- Atrial (see *Auricle* or *Auricula*)
- Atrial septal defect 213-20
- angiography 220
 - arachnodactyly in 216
 - bacterial endocarditis in 220
 - cardiac catheterisation 218
 - clinical features 216
 - electrocardiogram 218
 - embryology 213
 - hemodynamics 216
 - incidence 216
 - mitral stenosis in 216
 - prognosis 220
 - treatment 220
 - X ray 52 218
- Atrophy of the heart 38
- Atropine
- and cholinergic drugs 138
 - Atropine (contd.)
 - in digitalis poisoning 148
 - in heart block 113 118 271
 - insensitivity (in myxœdema) 496
 - intravenous dose (maximum) 118
 - in nodal rhythm 117
 - in paroxysmal cardiac dyspnoea 190
 - in pulmonary embolism 449
 - in sino auricular block 115
 - syncope 197
 - vagal paralysis from 110
- Aural vertigo 200
- Aureomycin 339
- Auricle left (and see *Left auricular*)
- aneurysmal dilatation 50 52 284
 - appendage 26 216
 - dilatation 47-52 288
 - expansile pulsation 283
 - hypertrophy 285
 - and left bronchus 93
 - in left ventricular failure 47
 - in Lutembacher's syndrome 218
 - in mitral incompetence 283
 - in mitral stenosis 47 288
 - and œsophagus 47
 - X ray 26 47
- Auricle right (and see *Right auricular*)
- dilation 59
 - in right ventricular failure 175
 - in tricuspid incompetence 304
 - in tricuspid stenosis 307
 - X ray 23 7 59
- Auricular ectopics 127
- Auricular fibrillation 145-50
- age and 147
 - angina pectoris and 147
 - in aortic valve disease 147
 - in atrial septal defect 218
 - in bacterial endocarditis 147 331
 - circus movement 145
 - clinical features 147
 - in congenital heart disease 147
 - in congestive failure 147
 - diagnosis 147
 - in digitalis intoxication 3 2
 - digitalis therapy 147
 - in diphtheria 313
 - in electric shock 531
 - electrocardiogram 145
 - embolism in 150
 - etiology 147
 - E waves 145
 - in head injuries 531
 - heart block in 145
 - hypertensive 424
 - incidence 147
 - in infections 147 149
 - jugular phlebogram 19 170
 - 1 contractions 145
 - VI contractions 145

- Auricular fibrillation (contd.)
 mechanism 145
 in Meniere's syndrome 531
 in meningitis 531
 in mitral stenosis 233
 in myocardial infarction 383
 in myocarditis 147 312 317
 in normal hearts 147
 paroxysmal 147 487
 physiology 145
 in Pick's disease 147 349
 in pneumonia 147
 in pregnancy 149 509
 prognosis 147
 in pulmonary heart disease 147 462
 quinidine therapy 148
 in rheumatic carditis 273
 syncope in 147
 thyrotoxic 487 489 492
 traumatic 531
 treatment 147
 in tricuspid valve disease 304 307
 vagal 113 531
 venous pulse 19 170
 in Wolff Parkinson White syndrome 141
- Auricular flutter 141-5
 angina pectoris in 142 408
 circus movement 141
 clinical features 142
 in congestive failure 142
 diagnosis 142
 digitalis therapy 144
 electrocardiogram 142
 etiology 142
 of waves 142
 first heart sound 142
 heart block in 142
 hypertensive 425
 incidence 142
 in infections 142
 in isolated myocarditis 321
 L contract ions 145
 M contract ions 145
 mechanism 142
 in mitral stenosis 293
 in myocardial infarction 389
 in myocarditis 317
 in normal hearts 14
 physiology 142
 prognosis 145
 in pulmonary heart disease 144 462
 quinidine therapy 145
 tumult 531
 treatment 144-5
 vagal 113 531
- Auricular gallop 164
 Auricular leads 73
 Auricular pressure (normal) 213
 Auricular pulse wave (see a wave)
 Auricular septum 213
- Auricular sounds 163 4
 in gallop rhythm 164-5
 in heart block 120
 Auricular tachycardia 132-4
 Auricular T wave (Ta wave) 8
 Auriculoventricular block (see Heart block) 117-24
 Auriculoventricular node 108
 Auscultation 7 10
 Auscultatory gap 4
 Austin Flint murmur 296
 Autonomic imbalance 24 35
 Autonomic nervous system (see also) 110
 Axis deviation 85
 Ayerza's disease 474
- Babcock's operation 363
 Back pressure theory 154
 Backward tilting of the apex 462
 Bacterial endarteritis 330
 in coarctation of the aorta 211
 in patent ductus 227
 Bacterial endocarditis 330-7
 acute 330
 anaemia in 331 332
 in aortic incompetence 291 330
 in aortic stenosis 303 330
 in atrial septal defect 220
 in auricular fibrillation 331
 bacteria free stage 33
 bacteriology 330
 in bicuspid aortic valve 12
 blood culture 335
 Bracht-Wachter bodies 331
 clinical features 331
 clubbing 333
 in coarctation of the aorta 211
 in congenital heart disease 330
 course 335
 diagnosis 335
 differential diagnosis 506
 embolic nephritis 334
 embolism 333
 etiology 330
 in Fallot's tetralogy 243 244
 fever 331
 haematuria 334
 heparin therapy 336
 incidence 331
 leucocyte count 22 332
 malignant 330
 in mitral valve disease 294
 myocarditis 331
 mycotic aneurysm 333
 nephritis 334
 Ole's nodes 333
 in patent ductus 227-9
 pathology 330
 pneumothorax 337-8
 proformation of aortic sinus 331

- Bacterial endocarditis** (contd)
 petechiæ 332
 prognosis 335
 in pregnancy 510
 pulmonary embolism 333 445
 in pulmonary stenosis 231 243
 renal lesions 334
 retinopathy 334
 in rheumatic heart disease 294 330
 splenomegaly 332
 subacute 330
 subarachnoid hæmorrhage 337
 sulphonamides in 336
 in syphilitic aortic incompetence 330
 treatment 336-9
 uræmia 336
 urine 334
 in ventricular septal defect 223 330
- Bainbridge reflex** 110 112
 in arterio venous aneurysm 514
 in heart failure 110 178
- Ball valve thrombus** 193
- Ballistocardiography** 21
- Bandaging the lumba** 311 200
- Barium chloride** in heart block 124
 toxic effects of 326
- Barium swallow** 27
- Basal metabolic rate** 2- 488
 in heart failure 23
 in leukæmia 489
 in myxœdema 496
 in thyrotoxicosis 488
- Basal rates** 160
- Basal systolic murmur** 281-2 301-2
- Basophil tumour of pituitary** 436
- Bayley's triaxial reference system** 81
- Bazett's formula** 79
- Bed cardiac** 179
- Bed pan calling for** 449
- Bed rest** in angina pectoris 384
 causing plebothrombosis 446 454
 in diphtheria 313
 in heart failure 179
 in myocardial infarction 406
 in rheumatic carditis 275
- Beer drinker's heart** 326
- Bell stethoscope** 7 8 288
- Benign hypertension** (see *Hypertension*)
 416
- Benign pericarditis** 354
- Benzodioxan** 421
- Benzoic acid** 338
- Beri beri** 502 514-16
 adrenalin in 516
 alcohol and 514 515
 arteriovenous shunt in 516
 cardiac output 502 515
 clinical features 502 515
 diagnosis 516
 electrocardiogram 516
- Beri beri** (contd)
 etiology 514-15
 hæmodynamic 502 515
 incidence 515
 pathology 516
 peripheral neuritis 516
 physiology 515
 pitressin in 516
 in pregnancy 515
 in thyrotoxicosis 491 515
 treatment 516
 X ray 57
- Bernheim's syndrome** 425
- Berry aneurysm** 211
- Bicuspid aortic valve** 209 212
- Bigeminal pulse** 129 130
- Bilharzia** 461
- Bilirubin blood** (in congestive failure) 175
- Bing's test** 244
- Bipolar leads** 68
- Bistiens pulse** 300
- Bismuth** (in syphilis) 367
- Blalock Taussig operation** 243
- Blast injury** 355 429
- B L B mask** 435
- Blood calcium** effect on electrocardiogram
 79 326
- Blood cholesterol**
 in ischemic heart disease 373 381 385
 497
 in myxœdema 496 497
 in thyrotoxicosis 488
- Blood count** 22
 in heart failure 175
- Blood culture** 335
- Blood gas analysis** 16
- Blood potas sum** (see *Potassium*) 326
- Blood pre sure** (and see *Hypertension*) 3-5
 in angina pectoris 374 375 429
 in aortic incompetence 295 297
 in aortic stenosis 301
 in arteriovenous aneurysm 514
 in atherosclerosis 5
 in auricular fibrillation 4
 auscultatory lap 4
 basal 5 416
 in cardiac compression (tamponade) 344
 345 348
 and cardiac output 178
 casual 5 416
 clinical assessment 3
 in coarctation of the aorta 208 211
 in circulatory failure 314
 in congestive failure 178
 in constrictive pericarditis 349
 in coronary thrombosis 387
 diastolic (normal) 5
 difference in two arms 5
 in ectopic beats 4
 in essential hypertension 422

- Blood pressure (contd)
 in heart block 122
 high (see *Hypertension*) 416-40
 in left ventricular failure 160
 in legs 5 208
 low (see *Hypotension*) 5 195 196 311 314
 lowering agents 381 437 439
 mean 5
 measurement 3-5 416
 in myocardial infarction 389
 in myxœdema 497
 normal 5
 in obesity 416
 in paroxysmal hypertension 421
 in patent ductus 223
 in pericardial effusion 344
 physiology 420
 in Pick's disease 349
 posture and 5
 in psychoneurosis 540
 in pulmonary embolism 448 449
 in pulmonary heart disease 465
 in shock 311
 systolic (normal) 5
 in syncope 194-7
 in thyrotoxicosis 494
 waxing and waning 314
 Blood proteins (in œdema) 173
 Blood sedimentation rate (see *E S R*) 22
 Blood urea in heart failure 22
 Blood viscosity (in hypertension) 420
 Blood volume
 in acute nephritis 171 327
 in anæmia 173 503
 in congestive failure 173
 in nephritic œdema 173
 Bloodless venesection 195
 Blows over the heart 528 32
 Blue baby 204 236
 Bohn's sign 223
 Bone decalcification of (in thyrotoxicosis) 488
 blood flow (in Paget's disease) 517
 Bone marrow culture 335
 Boot shaped heart 429
 Bowles stethoscope 8
 Brachial arteries examination of 3
 Bracht-Wachter bodies 331
 Bradycardia (see *Sinus bradycardia*) 112-124
 atropine effect on 113 197
 and auricular fibrillation 113
 cardiac enlargement in 35 113
 in heart block 140
 in sino auricular block 114-15
 Brain (see *Cerebral*)
 Brandy (see *Alcohol*)
 Brønham's sign 514
 B cath holding tests 418 536
 Breathlessness (see *Dyspnoea*) 158-60 536
 Bright's disease (see *Hypertension* and *Chronic nephritis*) 418 422-40
 Broad diameter 34
 Broadbent's sign 351 352
 Brock's operation
 for Fallot's tetralogy 244 306
 for mitral stenosis 308
 for pulmonary stenosis 231
 Bromides 130 437 491
 Bronchial arteries
 in Fallot's tetralogy 236
 hæmorrhage from 434
 in pulmonary atresia 250
 Bronchial asthma 464 465
 treatment 473
 Bronchial compression
 from aneurysm 360
 from left auricular enlargement 293
 from pericardial effusion 344
 Bronchial spasm 190 473
 Bronchial veins 161
 Bronchiectasis 464
 Bronchitis and emphysema 464 465
 treatment 473
 Bronchitis in mitral stenosis 293
 Broncho-pulmonary anastomosis 236 250
 Bruit (see *Murmur*)
 Bruit de gallop 166
 Bruit de Roger 22
 Buerger's disease 372
 Bulbus cordis 229
 Bullet in pulmonary artery 458
 in heart 525-7
 Bundle branches 108
 Bundle branch block 125-7
 alternating 126
 anatomy 125
 in atrial septal defect (right) 218
 clinical features 126
 diphtheritic 312
 electrocardiogram 90-3
 etiology 126
 experimental 125
 in familial card omegaly (left) 205
 functional 138 218 451
 gallop rhythm in 126
 histology 125
 in hypertension (left) 4 9
 kymography in 125
 left 90 1 6
 in mitral stenosis (right) 290
 in myocardial infarction 39 397
 in myocarditis 318
 nomenclature 125
 paroxysmal 126
 in paroxysmal tachycardia 128
 partial 126 218 90
 pathology 125

- Bundle branch block (contd)
 - physiological 139
 - prognosis 127
 - in pulmonary embolism (right) 431
 - in pulmonary heart disease (right) 467
 - right 93 126
 - split second sound in (right) 10 126 218
 - in syphilitic aortic incompetence (left) 363
 - transient 126 451
- Bundle of His 108
- branches of 141
- calcification of 120 517
- fibrosis of 120
- haemorrhage into 120 531
- Bundle of Kent 141
- Burns 195
- C wave of phlebogram 19 170
- Cachexia - cardiac 174
- Cæsarian section 510
- Caffeine as diuretic 185
- Calcific aortic stenosis 302-3
- Calcification of (or in)
 - aneurysm 363
 - aorta 38 30 358 365 174
 - aortic valve 61 301 517
 - arteries 374 517
 - coronary arteries 374 397
 - goitre 485
 - mitral a/c 63 284 517
 - Paget's disease 517
 - pericardium 61 349 352
- Calcium (blood)
 - effect on Q-T interval 104
 - gluconate for circulation time 13
 - lactate in aneurysm 363
- Campbell de Morgan's spots 332
- Cannon waves 118 120 170
- Capillary
 - blood pressure 462
 - electrometer 65
 - fragility test 332
 - permeability in oedema 173
 - pressure pulmonary 462
- Capillary pulsation 3 297
 - in aortic incompetence 297
 - in arteriovenous aneurysm 514
 - in cor pulmonale 465
 - in hyperkinetic circulatory states 502
 - in thyrotoxicosis 487
- Capillary resistance test 332
- Carbo amino acetyl choline (carbochol doryl) 138 200
- Carbon dioxide
 - and cerebral blood flow 200
 - in Cheyne Stokes respirations 167
 - in hyperventilation 536
 - retention (in emphysema) 464
- Carbon monoxide poisoning 102 468
- Carcinoma
 - embolic 458
 - pericardial 354
 - phlebothrombosis in 446
- Cardiac accelerator nerves 109
- Cardiac aneurysm 404 405
 - electrocardiogram 391 404
 - traumatic 529
 - X-ray 44 397
- Cardiac asthma 160
- Cardiac atrophy 38
- Cardiac bed 179
- Cardiac cachexia 174
- Cardiac catheterisation (in) 13-17
 - atrial septal defect 218 234
 - Eisenmenger's complex 245
 - Fallot's tetralogy 242
 - heart failure 154
 - left ventricular failure 160
 - mitral stenosis 285 292
 - patent ductus 227
 - patent foramen ovale 213 234
 - pulmonary stenosis 231 232
 - transposition 246
 - ventricular septal defect 21
- Cardiac chair 179
- Cardiac cirrhosis 172
 - in Ick's disease 348
 - in tricuspid disease 306
- Cardiac compression (see *Cardiac tamponade*) 345 9
- syncope in 194
- Cardiac dilatation (see *Cardiac enlargement*)
- Cardiac displacement 35
- Cardiac dullness 7
 - in emphysema 465
 - in pericardial effusion 345
- Cardiac enlargement (general) 61
 - in anaemia 506
 - in arteriovenous aneurysm 514
 - in athletes 35
 - in bradycardia 35 113
 - congenital 205
 - diagnosis 321 348
 - in fever 266
 - in heart block 12
 - in heart failure 175
 - in isolated myocarditis 318
 - in myxoedema 497
 - in nutritional cardiopathies 3 1
 - in pregnancy 508
 - in rheumatic carditis 266
 - in sino auricular block 115
 - spurious 35 38
 - in thyrotoxicosis 487
 - in toxic myocarditis 317
- Cardiac failure (see *Congestive failure*) 154 191

- Cardiac impulse (see *Apeæ beat*) 6
 heaving 424
 hyperdynamic 222 424
 impalpable 216 465
 tapping 216 237 256
 thrusting 223 424
 tumultuous 218
 Cardiac infarction (see *Myocardial infarction*)
 Cardiac insufficiency (see *Congestive heart failure*) 154-91
 Cardiac measurements 34
 Cardiac negro 474
 Cardiac nerves 107 386
 Cardiac neurosis 535-45
 and anæmia 506
 differential diagnosis 542-3
 etiology 535
 psychiatric aspects 541
 and rheumatic fever 276
 signs 540
 symptoms 536-8
 and thyrotoxicosis 481 485
 treatment 543
 Cardiac output 15 154
 in anæmia 502 503
 in arteriovenous aneurysm 514
 in bradycardia 113 115
 in beri beri 515
 in congestive heart failure 154-7
 on exercise 16
 in heart block 122
 high 155 6 50
 in left ventricular failure 158-60
 low 154-6
 measurement of 15
 in mitral stenosis 285
 normal 16
 in Paget's disease 517
 in pericardial effusion 344-5
 physiology 110 154 493
 in Pick's disease 349
 in pregnancy 507
 in pulmonary embolism 447 456
 in pulmonary heart disease 464
 in syncope 193-7
 in tachycardia 110-12 132
 in thyrotoxicosis 480 489
 Cardiac rotation (see *Rotation*)
 Cardiac rupture 404 528
 Cardiac standstill 115 193
 Cardiac syncope (see *Syncope*) 193-4
 Cardiac tamponade 345-8
 from direct injury 355 526 5 7
 from dissecting aneurysm 5
 from hæmopericardium 355
 from indirect injury 329
 in malignant pericarditis 354
 from perforated infarct 404
 and syncope 194
 Cardiac tamponade (contd.)
 in tuberculous pericarditis 353
 Cardiac vector 80
 Cardiolytic 351
 Cardio omentopexy 386
 Cardio thoracic ratio 34
 Carditis (see *Myocarditis*)
 rheumatic 255-7
 toxic and other form 311 28
 Carotophylline (see *Iminophyllin*)
 Carey Croombs murmur 270
 Caronamide 338
 Carotid artery
 examination of 3
 kinking of 423
 tortuosity of 422
 Carotid compression (bilateral) 201
 Carotid pulsation 169
 Carotid sinus pressure 138
 in auricular flutter 133 142 144
 and cardiac standstill 115 196
 in heart block 123
 in paroxysmal tachycardia 132-3 137 138
 physiology 196-7
 in sinus tachycardia 111
 Carotid sinus syncope 196-7
 Cations 65
 Cellophane treatment of aneurysm 364
 Cerebral anoxia 201
 Cerebral blood flow 196 200-1
 in hypertension 422
 in hyperventilation 200 526 529
 in pulmonary heart disease 465
 in syncope 193-6
 Cerebral embolism 16 243 333
 Cerebral hæmorrhage 433
 Cerebral symptoms in heart failure 174
 in hypertension 422
 Cerebral syncope 197 200
 Cerebral thrombosis 434
 Cervical sympathetic 386
 Chagas disease 315
 Chl cardiac 179
 Chemotherapy in bacterial endocarditis 336
 in syphilitic aortitis 367
 Chest leads normal 68-73
 Cheyne Stokes respiration 167 190
 Chl d b rth pulmonary embolism in 446
 Chloroform 150 326
 Chloromycetin 339
 Cholecystitis 405
 Cholesterol (see *Blood cholesterol*) 373 496
 Cholergic drugs action of 138 199-200 516
 in paroxysmal tachycardia 138
 in syncope 199
 Chordæ tendineæ 28
 rupture of 530

- Chorea 461
treatment 475
- Chorion epithelioma 459
- Chronic bronchitis 464 473
- Chronic constrictive pericarditis (see *Pick's disease*) 348-51
- Chronic nephritis (see *Hypertension*) 417-418 423 425 433
- Cinematography 31
- Circoid aneurysm congenital 511
- Circulatory failure
and coronary insufficiency 408 10
in diphtheria 311
in infections 314
in pulmonary heart disease 465
and syncope 194-6
- Circulation time 12-13 31 161
in beri beri 515
in hyperkinetic circulatory states 502 515
in left ventricular failure 163 425
in mitral stenosis 292
in myxoedema 496
in right to left shunt 243
in right ventricular failure 163
in thyrotoxicosis 489
- Circus morient
in auricular fibrillation 145
in auricular flutter 141-144-5
in paroxysmal tachycardia 141
- Cirrhosis (and see *Cardiac cirrhosis*)
effect on the circulation 517
in haemochromatosis 321
and nutritional cardiopathy 321
and xerema 172
- Clinical examination 2-11
- Chimeric (see *Mosaicism*)
- Clockwise rotation (about longitudinal axis) 70 82 85
in Fallot's tetralogy 242
in hypertension 426
in pulmonary heart disease 465
in right ventricular enlargement 89
- Clotting (see *Thrombosis etc*) 444-6
- Clotting time 454
- Cloudy swelling 313
- Clubbing of fingers 333
in aneurysm 360
in bacterial endocarditis 333
in congenital heart disease 237
mechanism 333
in pulmonary heart disease 462 465
- Coarctation of the aorta 206-12
associated phenomena 209
blood pressure 208
classification 207
clinical features 208
collateral circulation 207
complications 208 209 211
embryology 206
haemodynamics 207
Coarctation of the aorta (contd)
prognosis 211
treatment 211
X ray 39 208-9
- Cœur en sabot 57 237 240
- Cold in angina pectoris 429
- Cold drinks 138
- Cold extremities 177 485
- Cold hypersensitivity to 496
- Cold pressor test 418
- Collapse of lung
in aneurysm 360
in kyphoscoliosis 474
in mitral stenosis 293
in pericardial effusion 344
in rheumatic fever 266
X ray 35
- Collapsing pulse (see *Water hammer*) 29,
- Colloid 478
- Coma 201
cardiac 434
in dissecting aneurysm 523
- Compensation neurosis 532 644
- Compensatory pause 127 129
- Complemental air 17
- Complete heart block (see *Heart block*) 120-4
- Concato's disease 333
- Concordant left ventricular preponderance 88 301 429
- Conducting system anatomy 108 125
physiology 108-9
- Congenital aortic stenosis 212 302
circoid aneurysm 511-13
cystic lung 464
dilatation of the pulmonary artery 212
heart block 120
heart disease (see *Anomalous regurgitation*) 203 250
classification 203-4
etiology 203
incidence 203
hypertrophy of the heart 205
- Congestive heart failure (and see *Left and right ventricular failure*) 154-91
acropuncture 189
in acute nephritis 327
albuminuria 21
in anaemia 503
in aortic stenosis 303
in arteriovenous aneurysm 514
ascites 174
in auricular fibrillation 147
in auricular flutter 142
back pressure theory 154
in bacterial endocarditis 331 335
basal metabolic rate 23
in beri beri 515
blood pressure 178
blood urea 22

Congestive heart failure (contd.)

- cachexia 14
- in cardiac infarction 403 407
- cardiac output 154-6
- cerebral symptoms 174
- circulation time 163
- clinical features 3 168-78
- compensated 155
- cyanosis 177
- definition 154-6
- diet 179 185
- digitalis therapy 179-84
- embolism 444-6
- erythrocyte sedimentation rate 22 175
- etiology 156-7 168
- gallop rhythm 165
- in heart block 122 124
- heart sounds 1, 8
- hepatic distension 171
- hydropneumothorax 174
- hydrothorax 174
- in hypertension 425 435
- insomnia 179
- jaundice 175
- liver 171
- low sodium diet in 185-9
- low voltage electrocardiogram 164
- mechanism 154-6
- mercurial diuretics in 184
- in mitral incompetence 284
- in mitral stenosis 168 285 293
- in myocarditis 312 317 318
- in myxoedema 496
- oedema 172-3
- in Paget's disease 517
- in paroxysmal tachycardia 132
- polycythæmia 22 175
- in pregnancy 509
- prognosis 178-9
- pulmonary embolism in 175 444-6
- in pulmonary heart disease 449 462 473
- pulse rate 178
- renal blood flow 173
- renal function tests 22 173
- reticulocytæmia 22
- in rheumatic carditis 270 275
- sodium retention 173
- staphylococci in 184
- in syphilitic aortitis 365
- tachycardia 110
- thiouracil in 190-1
- thrombosis in 444-6
- in thyrotoxicosis 487 493 495
- treatment 179-91
- urine 174
- vasoconstriction 178 285 447
- venocut in 179
- venous pressure 154-6 168-71
- venous thrombosis in 175 445 446
- X-ray appearances 59 175

Constrictive pericarditis (see *Pick's disease*)

348-51

Continuous murmur

- in arteriovenous aneurysm 513 514
- artificial 24
- broncho-pulmonary 229 236 250
- in patent ductus 223
- in perforated aortic sinus 229 524
- in thyrotoxicosis 486
- in thiouracil goitre 493
- venous 22

Contusion of the heart 52

Conus lifting 216 222 462

Conus of right ventricle 27 47

Convulsions in

- cardiac syncope 193
- epilepsy 198
- Stokes-Adams attacks 123
- vaso-motor syncope 198

Cor pulmonale (see *Pulmonary embolism* and *Pulmonary heart disease*) 444 461

Coramine 190 455

Coronary angitis 372 374

rheumatic 257 294

Coronary arteries

- anatomy 387
- anomalous 206
- atherosclerosis 372-4
- calcification 374 397
- dissection 522
- in hypertensive heart disease 375 433
- injury 526 530 532
- subintimal hæmorrhage 374 384
- in syphilitic aortitis 366

Coronary blood flow 366 433

Coronary embolism 372 374

Coronary insufficiency acute 408-10

Coronary occlusion 370 372-4 380 386 388

Coronary ostia stenosis of 366

Coronary sinus 108

Coronary sinus rhythm 116 136

Coronary spasm 377 386

Coronary T wave 93-7 389-97

Coronary thrombosis (see *Myocardial infarction*) 380 386

definition 387

etiology 374

site of 387 8

spreading 384

treatment 384

Coronary thromboangitis 372 374

Coronary vein ligation of 380

Corrigenda 297

Cortisone 275

Coupled beats 19 130

Crush injuries 355 529

Current of injury 94 99

Cushing's syndrome 436

- Cyanide 12
 Cyanosis 177
 in mitral stenosis 285
 Cystic medial necrosis 521

 D A H (see *Cardiac neurosis*) 535
 Da Costa's syndrome (see *Cardiac neurosis*)
 535-45
 Dactylic metre 164
 Dalrymple's sign 481
 Death (see *Sudden death*)
 Decholin regulation time 12 31
 Decompensation (see *Congestive heart failure*) 554-91
 Dementia
 in heart failure 174
 in hypertension 422
 Depolarized cell 65
 Depressed sternum 35
 Depressor reflexes 113 196
 Development of
 aortic system 206
 auricular septum 213
 heart 203
 pulmonary arteries 206
 Dextrocardia 204
 Dextroposition of aorta 236
 Diabetes mellitus
 in angina pectoris 371 378
 in atherosclerosis 373
 in ischaemic heart disease 408
 Diabetic retinopathy 424
 Diameters of the heart 34-5
 Diaphragmatic hernia 377 406
 Diastase urinary 406
 Diastolic blood pressure (see *Blood pressure*) 5
 Diastolic murmurs (see *Aortic Mitral etc.*)
 Diastolic rebound 351 352
 Diastolic report 349
 Diastolic shock 349
 Diastolic thrills (see *Aortic Mitral etc.*)
 Dicoumarol in
 angina pectoris 384
 cardiac infarction 406 7
 pulmonary embolism 451
 thrombo embolic states 454
 Dirotic pulse in circulatory failure 314
 Dirotic wave (of the pulse) 20
 Diet
 in heart failure 179 185
 in hypertension 438
 low sodium 185-9
 in myocardial infarction 406
 Digital throbbing (see *Vasodilatation*) 502
 Digitalis
 action 144 179 84
 administration 144 147 148
 atropine antagonism 148 325
 in auricular fibrillation 147-8
 Digitalis (contd.)
 auricular fibrillation from 322
 in auricular flutter 144
 and clotting time 446
 in congestive failure 179 84
 contra indications 124 190 313 317
 407
 in diphtheritic carditis 313
 dose 144 147-8 184
 ectopic beats from 130 322
 electrocardiogram 101 325
 in heart block 124
 heart block from 322
 in left ventricular failure 179-84 190
 in myocardial infarction 407
 nodal rhythm from 3 2
 in paroxysmal tachycardia 138
 paroxysmal tachycardia from 322
 preparations 148
 in pulmonary heart disease 464 474
 and Q T interval 325
 in rheumatic carditis 275
 sudden death from 322
 T wave 101
 toxic effects 148 322-5
 in toxic myocarditis 317
 ventricular fibrillation 150 322
 Digitoxin (see *Digitalis*) 148
 Digoxin (see *Digitalis*) 148
 Diiodotyrosine 478
 Diodone 31
 Diodrast 31
 Diphtheria
 circulatory collapse in 311
 myocarditis in 311-13
 Dipole 65
 Dipping (on to liver) 172
 Disordered action of the heart (see *Cardiac neurosis*) 535
 Dissecting aneurysm of the aorta 521-3
 haemopericardium in 355
 Disseminated lupus 315
 Dissociation auriculo ventricular (see *Heart block*) 120-4
 Distortion by diverging X rays 29 38
 Diuretics 184-5
 Diuretin 185
 Dizziness 197-201 536 538
 Dock's sign (rib notching) 208
 Dominance
 left ventricular 87-8
 right ventricular 88-9
 Dorsal artery of the foot
 emboli in 333
 palpation of 3 333
 Doryl (see *Cholinergic drugs*) 138 200
 Double aortic knuckle 209
 Double layer 65
 Doublet 65
 Dresden clinalock 300

- Drinking (see *Alcohol*) 326
 Dropped beats
 in partial heart block 119
 in sino-auricular block 114
 Dropsy (see *oedema*) 172-3
 Ductus arteriosus 223-9
 development of 206
 Dullness cardiac 7
 in emphysema 465
 in pericardial effusion 345
 Duroz er's sign 297
 Dysenteric polyarthritis 260
 Dysphagia (from)
 aneurysm of the aorta 361
 goitre 486
 unfused aorta 429
 Dyspnoea (in)
 heart failure 158-60
 mitral stenosis 285
 psychoneurosis 536
 Ectopic beats 127-31
 auricular 127 133 342
 diagnosis 129
 from digitalis 322
 in diphtheria 312
 etiology 130
 experimental 130
 interpolated 131
 in myocardial infarction 389 397
 nodal 128
 in pregnancy 507
 treatment 130
 ventricular 129
 Effort syndrome (see *Cardiac neurosis*) 535
 Effort tolerance test (in)
 angina pectoris 378
 limited cardiac reserve 171
 psychoneurosis 540
 Einthoven's string galvanometer 67
 Einthoven's theory 68 77
 Einthoven's triangle 77 81 86
 Esmarch's complex 244-f
 X-ray 52
 Electric shock 531
 Electrical alternation 167
 Electrical axis 81 85
 deviation of 85-7
 fixation of 351 352
 Electrocardiogram (and see *Electrocardiography*)
 acute cor pulmonale 449-51
 acute coronary insufficiency 410
 acute nephritis 327
 anæmia 504
 angina pectoris 101 378 382
 anoxic states 10 410
 aortic incompetence 298
 aortic stenosis 301
 asphyxia 101 410 467
 electrocardiogram (contd.)
 atrial septal defect 218
 auricular ectopic beats 12
 auricular f waves 142 144 145
 auricular fibrillation 145
 auricular flutter 142
 auricular standstill 14f
 auricular tachycardia paroxysmal 13 -
 135
 auricular T waves 8
 axis deviation of 85
 beriberi 516
 bundle branch block (see *BBB*)
 left 90
 right 93
 carbon monoxide poisoning 102 411
 cardiac injuries 529
 cardiac standstill 115
 carditis 104 312 317 318
 circulatory failure 102 410
 clockwise rotation 70 82 85
 concordant left ventricular preponderance 88 301 429
 congenital hypertrophy of the heart 205
 congestive failure recurrent 104
 constrictive pericarditis 349
 cor pulmonale 449 465
 coronary thrombosis 93 389
 current of injury 94
 dextrocardia 204
 digitalis effect 101 325
 diphtheritic carditis 104 312
 dissecting aneurysm 52-
 ectopic beats 127-31
 extra systoles (interpolated) 131
 f waves 142-5
 Fallot's tetralogy 242
 familial cardiomegaly 205
 Fiedler's carditis 318
 hæmorrhage 102 410
 heart block
 complete 120
 partial 118
 retrograde 117
 horizontal heart 77
 hypertensive heart disease 87 4 6
 hypocæmia 79 326
 hypoxia 101 410 467
 idiopathic dilatation of pulmonary artery 212
 ischemic heart disease 93 101 3 8
 382
 isolated myocarditis 318
 left auricular enlargement 83
 left axis deviation 86
 left ventricular enlargement 87
 left ventricular failure 83
 left ventricular preponderance 87 4 6
 long P-R interval 118
 low voltage curves 103

Electrocardiogram (contd)

- mitral incompetence 283
- mitral P wave 83 290
- mitral stenosis 83 290-2
- myocardial bruising 529
- myocardial infarction 93-7 389-97
- myocarditis 104 312 317 318
- myxoedema 103 494
- nodal ectopic beats 1 8
- nodal rhythm 117
- normal 70-9
- P mitrale (see *P*) 83 290
- P pulmonale (see *P*) 83 170 467
- P R interval (see *P R*) 78
- P wave (see *P*) 67 78 83 467
- Pardee T wave 97 391
- paroxysmal tachycardia 132-41
 - auricular 132-5
 - nodal 136
 - ventricular 137
- patent ductus 224
- pericarditis 99 342-4
- Pick's disease 349
- posture effect of 86-7
- potassium effect 104 326
- pregnancy 507
- pulmonary embolism 449-51
- pulmonary heart disease 463 465-70
- pulmonary stenosis 231 242
- Q wave (see under *Q*) 69 78 97
- QRS complex (see under *QRS*) 68-70 85-93
- QRS patterns basic 73
- Q T interval (see *Q Tc*) 79
- R wave (see *QRS complex*)
- RS T segment (see under *RS T*) 79 93-105
- rheumatic carditis 269
- right axis deviation 87
- right entricular dominance 88
- rotation of the heart 70 77 82 85
- S wave (see *QRS*) 69 79
- S T interval (see *RS T*) 79 93-105
- sino auricular block 144
- sinus arrhythmia 109
- sinus bradycardia 113
- T wave (see under *T*) 67 73-4 93-105
- Ta wave 67
- thyrotoxicosis 487 490
- toxic myocarditis 317
- tricuspid atresia 250
- tricuspid stenosis 307
- U wave (see under *U*) 79-80
- uræmia 104 326
- vaso motor syncope 101
- ventricular ectopic beats 129
- ventricular escape 114
- ventricular fibrillation 150
- ventricular septal defect 222
- vertical heart 77

Electrocardiography (and see *Leads*) 65-151

- accession wave 68
- basic patterns 74
- cardiac vector 80
- central terminal 68
- chest lead 68-73
- dipole theory 65
- electrodes 68
- equations 75 77 78
- excitatory impulse 65
- history 65
- instruments 65 67
- limb lead 74-8
- paste for electrodes 68
- polarity 68
- principles 65-77
- regression wave 67
- standard lead 77
- standardisation 68
- technique 68 70 75
- theories 65-7
- unipolar lead 68
- Wilson's neutral electrode 68
- Electrokymography 29
- Elephant heart rate of 110
- Elongated heart 35
- Embolectomy pulmonary 455
- Embolie nephritis 334
- Embolism (see *Pulmonary embolism*)
 - air 457
 - in auricular fibrillation 150
 - in bacterial endocarditis 333
 - cerebral 404
 - in congestive failure 446
 - coronary 372
 - fat 457
 - foreign body 458
 - incidence 444
 - in isolated myocarditis 321 445
 - malignant 458
 - in mitral stenosis 493
 - in myocardial infarction 403-4
 - paradoxical 457
 - renal 334
 - splenic 332
- Emotion effect on
 - blood pressure 5 537 540
 - breathing 536 538
 - cardiac output 537
 - pulse rate 110 537 540
- Emphysema
 - clinical features 465
 - intrathoracic pressure 472
 - in kyphoscoliosis 474
 - lung volume 19 472
 - in pulmonary heart disease 464
 - residual air 19 472
 - right auricular pressure 472
 - treatment 473

- Emphysema (contd)
 vital capacity 19 464 472
 X ray 472
- Encephalopathy hypertensive 434
- Endocarditis
 bacterial 330
 rheumatic 255 280
 syphilitic 364
- Ephedrine
 in asthma 473
 in heart block 124
- Ephynal (vitamin E) 384
- Epilepsy 197
- Epinephrine (see *Adrenalin*)
- Epistaxis (in)
 coarctation of the aorta 208
 hypertension 434
 rheumatic fever 257
- Erosion (of)
 ribs 208
 vertebrae 361 363
- Erythema
 marginatum 264
 multiforme 264
 nodosum 264
- Erythrocyte sedimentation rate (in) 22
 congestive failure 175
 hypertension 22
 myocardial infarction 389
 pulmonary infarction 453
 rheumatic carditis 259
 syphilitic aortitis 365
- Erythrogenic toxins 256
- Essential hypertension (see *Hypertension*)
 416-39
- Ether circulation time 13 163
- Eupaverine 455
- Euphyllin (see *Aminophylline*) 190
- Ewart's sign 345
- Examination clinical 2-11
- Excitatory impulse 65
- Exercise (see *Effort tests*) 502
- Exhaustion 537
- Exophthalmic goitre (see *Thyrotoxicosis*)
 485
- Exophthalmic ophthalmoplegia 481
- Exophthalmos 481
- Expansile pulsation of
 aneurysm 38 363
 left ucle 52 283
 pulmonary artery 52 218
- Exploring electrode 68
- Extremities murmurs 281
- Extracardiac sound 165
- Extrasystoles (see *Ectopic beats*) 127-31
 interpolated only 31
- Eyes in thyrotoxicosis 481
- Eye pressure 111 138
- pp
- f waves
 in auricular fibrillation 146
 in auricular flutter 143
- Failure (see *Congestive heart failure*) 154-
 191
 circulatory 194 6 311 314
 congestive 154-91
 forward 154
 high output 156
 ischaemic 370
 left ventricular 157
 low output 156
 right ventricular 168
 syncopal 193
- Fainting (see *Syncope*) 194
- Fallot's tetralogy 236-44
 angiocardigram 242
 catheter findings 242
 circulation time 243
 clinical features 237
 complications 243
 definition 236
 electrocardiogram 242
 haemodynamics 236
 prognosis 243
 treatment 243
 X ray 57 240
- Familial cardiomegaly 205
- Fat apical or triangular pad 35
- Fat embolism 457
- Fat metabolism in atherosclerosis 373
- Fatigue 537
- Fatty degeneration of the heart 313
- Femoral arteriovenous aneurysm 513
- Femoral blood pressure 5 208
- Femoral phlebothrombosis (see *Phlebotrombosis*) 445 453
- Femoral vein ligation of 455
- Fever 502
 in naemia 506
 in bacterial endocarditis 331
 in dissecting aneurysm 52
 in myocardial infarction 389
 in myocarditis 316
 penicillin 338
 in pericarditis 341
 in pulmonary infarction 453
 in rheumatic carditis 259
 and sinus tachycardia 111
 thiouracil 385
- Fibrinous pericarditis 341
- Fibrosis of the heart
 scheme 388
 in isolated myocarditis 318
 nutritional 311
 humoral 307
 thyrotoxic 480
- Fick principle 15
- Fiedler's carditis 318-21
- Fingert clubbing (see *Clinical*) 333

- First heart sound
 in auricular flutter 142
 in bundle branch block 16
 in heart block 122
 in hyperkinetic circulatory state 10
 mechanism 9
 in mitral stenosis 286
 and P R interval 10
 weak 178
- First oblique position 20
- Fits
 epileptic 197
 Stokes Adams 122
- Fluids in congestive failure 179
- Fluorophotography 31
- Fluoroscopy
 technique 25-7
 value in
 aortic aneurysm 38 363
 atrial septal defect 52 418
 calcified valves 61 301
 mitral incompetence 52 83
 myocardial infarction 44 397
 myocardial foreign bodies 526
 pericardial effusion 61 345
 Pick's disease 61 349
- Fluores
 amyl nitrite 382
 cholinergic 138
 menopausal 200
 after Stokes Adams fits 121
 and syncope 200
- Fetal adenoma 486
- Fetal circulation 213
- Fetal endocarditis 203 229
- Fetal goitre 511
- Fetal rhythm 178
- Foramen ovale (see *Patent foramen ovale*)
 213
- Forearm blood flow
 in Cheyne Stokes breathing 168
 in thyrotoxicosis 489
- Foreign bodies
 embolism from 458
 intravascular 458
 myocardial 525 7
 pericardial 335 525-6
- Forced breathing 200 536
- Forward failure 154
- Fourth heart sound 163
- Fractures
 emboli from 446
 fat emboli from 457
- Frank's capsule 21
- Friction rub
 pericardial 341
 pleural 453
 pleuropericardial 342
- Fright 528
- Frustrated diet (in)
 heart failure 185
 hypertension 438
 myocardial infarction 406
- Functional
 aortic incompetence 424 503
 aortic systolic murmur 282
 arrhythmia 109
 bradycardia 112-17
 breathlessness 536
 bundle branch block 139
 dizziness 538
 fatigue 537
 headache 538
 hypertension 5 416
 mitral diastolic murmur 288
 mitral incompetence 282
 mitral systolic murmur 273 282
 pain 377 537
 palpitations 536
 parasternal murmur 220
 pulmonary incompetence (see under *Pulmonary incompetence*)
 signs and symptoms in anxiety states 540
 sweating 538
 syncope 194-201
 systolic murmur 281 503
 tachycardia 110-12
 tricuspid incompetence 303
- Functional ocular (in)
 aortic incompetence 298
 bacterial endocarditis 314
 diabetes mellitus 4 4
 examination of 5 423
 hypertension 4 3
 nephritis 424
 pulmonary heart disease 465
 rheumatic states 257
 superior vena cava obstruction 424
 fusiform aneurysm 38 353 364
- Gall stones 405
- Gallop rhythm (in) 163-f
 auricular 164
 bundle branch block 126
 congestive heart failure 163
 diphtheritic carditis 312
 hypertensive heart failure 165 4 4
 ischaemic heart failure 165 381
 isolated myocarditis 318
 left ventricular failure 158
 left ventricular stress 164 165 389 424
 long P R interval 164 165
 mitral stenosis 165
 normal persons 166
 nomenclature 166
 Pick's disease 165 349
 presystolic 164
 protodiastolic 165
 pulmonary heart disease 462 463

- Callip rhythm (contd.)
 rheumatic carditis 273
 right ventricular stress 164 462 465
 summation 165
 systolic 165
 tachycardia 164 165
 toxic myocarditis 2 3
 Calvanometer (tintha en s string) 67
 Cangelionectomy stellate 386
 Gas analysis 16
 Geiger counter 490
 German measles 203
 Giant *a* wave (see *a* trace) 170 229 306 462
 Gibson murmur 223
 Glycogen retention 205
 Glycosuria 22
 Goutre (see *Thyrotoxicosis*)
 adenomatous 479 486
 clinical features 480
 colloid 479
 diffuse hyperplastic 479 486
 endemic 479
 exophthalmic 479
 foetal adenomatous 486
 in foetus 495
 lymphadenoid 486
 malignant 486
 nodular 479
 simple 479
 substernal 485
 surgical treatment 490
 from thiouracil 493
 in thyroiditis 486
 Goldberger's augmented leads 75
 Goldblatt clamp 419
 Gonococcal
 endocarditis 330
 polyarthritides 260
 Gradient (ventricular) 82
 Graham Steell murmur 216 288
 Graves disease (see *thyrotoxicosis*) 478
 Great vessels development of 206
 Greter systolic click 165
 Growing pains 260
 Gumma of the heart 367
 Gunshot wound 354 525
 Haemangioma 511
 Haematemesis (in)
 aortic aneurysm 363
 hypertension 434
 Haematuria (in)
 bacterial endocarditis 334
 dicoumrol therapy 454
 dissecting aneurysm 53
 heart failure 334
 nephritis 334 433
 renal infarction 334
 Haemic murmur 503
 Haemochromatosis 321
 Haemoglobin
 in cyanosis 177
 general 22
 oxygen carrying capacity 177
 Haemolysis
 anti-streptococcal 257
 streptococcal 256
 Haemopericardium 354
 accidental 348
 in blast injury 529
 in dissecting aneurysm 522
 in gunshot wounds 525
 from indirect injury 529
 malignant 354
 in perforated cardiac infarction 404
 in ruptured heart 529
 in ruptured syphilitic aneurysm 363
 in stab wounds 527
 traumatic 354 529
 tuberculous 353
 Hemoptysis (in)
 atrial septal defect 216
 bacterial endocarditis 334
 congestive heart failure 175
 dissecting aneurysm 523
 hypertension 434
 isolated myocarditis 321
 left ventricular failure 160
 mitral stenosis 293
 pulmonary arteriovenous aneurysm 513
 pulmonary infarction 453
 ruptured aneurysm 363
 Hemorrhage
 in bundle of His 531
 capillary 257
 cerebral 433
 coronary 348
 and coronary insufficiency 408
 in dicoumrol poisoning 454 455
 effect on circulation 503
 electrocardiogram 102
 petechial 257 332
 purpuric 332
 retinal 334 423
 subarachnoid 434
 subintimal 374 384
 and syncope 19
 Hemothorax (in)
 dissecting aneurysm 523
 pulmonary infarction 453
 ruptured aneurysm 363
 Haldane's apparatus 16
 Hands (in)
 acrochordactyly 216
 cholesterol 261
 psoriasis 485
 thyrotoxicosis 485
 Hereditary rate 110
 Himotosis 486

- Hay type of dropped beats 119
 Headache
 anxiety 538
 in essential hypertension 422
 in fat embolism 458
 histamine 538
 in hypertensive encephalopathy 434
 in paroxysmal hypertension 421
 Trinitrin 538
 Head injury (and)
 auricular fibrillation 147 521
 auricular flutter 147
 sinus bradycardia 113
 Heart block (auriculo ventricular) 117-24
 in auricular fibrillation 147
 in auricular flutter 142 144
 auricular sounds 122
 blood pressure 122
 bundle branch block in 120
 cannon waves 118 122
 cardiac enlargement 35 122
 cardiac output 122
 clinical features 120
 complete 120-4
 congenital 120
 congestive failure 122 124
 digitalis 124 322
 diphtheritic 312
 electrocardiogram 118-21
 etiology 118 120
 first heart sound 122
 grades 117
 haemodynamics 122
 Hay type 119
 hypertensive 425
 ischaemic 389
 in isolated myocarditis 321
 in myocardial infarction 389
 in Paget's disease 517
 paroxysmal 120 122 123
 in paroxysmal tachycardia 123
 paroxysmal tachycardia in 123 124
 partial 118-20
 pathology 120
 prognosis 124
 pulse 120 122 124
 retrograde 117 128 129
 rheumatic 271 307
 Stokes Adams attacks 122
 sudden death 124
 syncope 120 193
 syphilitic 124 367
 in toxic myocarditis 317
 transient 118 120
 traumatic 531
 treatment 124
 cannon pulse 122 170
 ventricular fibrillation 123 124
 Wenckebach type 119
 \rav 122
 Heart dilatation of (see *Cardiac enlargement*)
 Heart disease
 etiology 1
 incidence 1
 mortality 1
 Heart enlargement etc. of (see *Cardiac enlargement etc.*)
 Heart failure (see *Congestive heart failure*)
 154-91
 Heart measurements 34
 Heart rate 110
 Heart shape 25-7 35
 Heart size 34 35
 Heart sounds 9 10 163 178
 auricular 120 163
 in congestive failure 178
 distant 178
 in emphysema 465
 extra 165
 faint 178 345 389 465
 first (see *First heart sound*) 9 10 163
 fourth 163 164
 in myocardial infarction 389
 in pericardial effusion 345
 reduplicated 164
 second (see *Second heart sound*) 10 163
 split 10 164
 third (see *Third heart sound*) 163-5
 weak 178
 Heart weight
 in anaemia 506
 in hypertension 425
 Helium 19
 Hemiplegia 16 318
 Heparin (in)
 acute coronary insufficiency 384
 administration of 454
 bacterial endocarditis 366
 cardiac catheterisation 16 242
 myocardial infarction 406
 pulmonary embolism 454
 rheumatic fever 266
 thrombo embolic states 454
 Hepatic cirrhosis (see *Cirrhosis*)
 Hepatic compression (manual) 171
 Hepatic distention 171-2
 failure 517
 function in thyrotoxicosis 488
 pain 171
 Hepatic pulsation
 pre systolic 229 306
 systolic 171 304
 Heredity (in)
 anxiety states 541
 congenital heart disease 203 20
 hypertension 418
 rheumatic fever 255
 Hering Breuer reflex 285
 Hernia diaphragmatic 377 406

- Herxheimer reaction 367 368
 High blood pressure (see *Hypertension*)
 416-40
 High output failure 156 302
 Hilar shadows 26
 His Bundle of (see *Bundle of His*) 108
 Histamine
 action of 200
 circulation time 12
 headache 538
 test for phaeochromocytoma 422
 History taking 1
 Hoarseness (in)
 aneurysm 361
 mitral stenosis 293
 myxoedema 496
 Homans sign 453
 Horizontal heart 35 77
 Horner's syndrome 361
 Hydræmia
 in acute nephritis 328
 in œdema 173
 venous pressure in 171
 Hydrogen for measuring lung volume 19
 Hydrogen peroxide for cleaning catheters
 17
 Hydropericardium 174 356
 Hydrostatic pressure 173
 Hydrothorax
 in congestive heart failure 174
 in left ventricular failure 44 161
 Hypercholesterolaemia
 in atherosclerosis 373
 in myxoedema 496-7
 Hyperpiesia (see *Hypertension*) 416-40
 Hyperpnoea (see *Hyperventilation*) 200 536
 Hyperkinetic circulatory states 155 502
 Anaemia 502
 arteriovenous aneurysm 513
 beriberi 514
 excessive 502
 fever 266 502
 hepatic failure 517
 Paget's disease 517
 pregnancy 507
 pulmonary heart disease 464 465
 thyrotoxicosis 487
 uræmia 321
 Hypertension 419 420
 Hypertensinase 419
 Hypertensinogen 419
 Hypertension (and see *Hypertensive heart disease*) 416-40
 albuminuria 433
 angina pectoris 375 429
 angiotonin 419
 atherosclerosis and 374 419 433
 auscultatory gap 4
 benign 423
 biochemical hypothesis 419-20
 Hypertension (contd)
 blood pressure 178 422 435
 blood viscosity 420
 blood volume 420
 cardiac output 420
 cerebral manifestations 433 434
 classification 416 417
 clinical features 421-34
 in coarctation of the aorta 208 418
 cold pressor test 418
 in coloured races 417
 course 434
 in Cushing's syndrome 436
 definition 416
 diastolic 416
 diet 437
 and emphysema 473
 epistaxis 434
 essential 417
 etiology 417 419
 experimental 417
 functional 416
 haematemesia 434
 hemoptysis 434
 heart in (see *Hypertensive heart disease*)
 424 435
 heredity and 418
 humoral theory 419-20
 incidence 417
 low sodium diet 437
 lumbo-dorsal sympathectomy 439
 malignant (see *Malignant hypertension*)
 419
 menopausal 416 437
 in mitral stenosis 417
 mortality 435 436
 nephrectomy for 436
 nephritic acute 327 417 424
 chronic 418 424 433
 nervous 5 420
 nocturia 433
 and obesity 416 436 437
 ocular fund 423
 papilloedema in 424
 paroxysmal 417 421
 pathogenesis 417
 persistent 422
 phaeochromocytoma and 421
 physiology 419-20
 polyuria 433
 in pregnancy 417 51
 prognosis 434-6
 pulmonary (see *Pulmonary hypertension*)
 461
 pulse 422
 pyelonephritis 418 436
 renal 417 420 436
 renal biopsy in 419
 renal function 420 433
 renal 419-20

- Hypertension (contd.)
 retinopathy 423-4 435
 sex incidence 417
 Smithwick operation 439
 stroke 433 436
 subarachnoid hæmorrhage 434
 surgical kidney 418
 surgical treatment 439
 sympathectomy 439
 symptoms 422
 systolic 416
 thiocyanates 438
 thyrotoxic 416
 transient 417 422
 treatment 436 40
 urine 433
 varieties 416
 vascular changes 419
 aortic constriction 420 422
 aneurysm 437
 Hypertensive encephalopathy 434
 Hypertensive heart disease (see *Hypertension*) 416
 angina pectoris in 429
 aortic incompetence in 424
 auricular fibrillation 424 425
 auricular flutter 425
 cardiac asthma 160
 cardiac enlargement 425
 congestive heart failure 42
 coronary circulation 433
 course 435
 electrocardiogram 87 4 6-9
 gallop rhythm 164 424
 heart block in 425
 left ventricular failure 157-63 425
 orthopnoea 158 425
 paroxysmal cardiac dyspnoea 160 45
 paroxysmal tachycardia in 45
 prognosis 435
 pulmonary oedema 160
 pulsus alternans 166 423 424
 \ ray 39 42 49 429
 Hypertensive heart failure 425 435
 Hypertensive retinopathy 423
 Hyperthyroidism (see *Thyrotoxicosis*) 478
 Hyperventilation
 in anxiety states 536
 in epilepsy 199
 and syncope 200-1
 Hypnotics 179
 Hypocalcæmia 79 326
 Hypoplasia of the aorta 42
 in atrial septal defect 218
 in coarctation 209
 in mitral stenosis 288
 Hypoplasia of the pulmonary artery 57 240
 Hypotension
 in cardiac tamponade 345
 in circulatory failure 311 314
 Hypotension (contd.)
 in isolated myocarditis 318
 in myocardial infarction 389
 orthostatic 195 200
 physiological 5
 in pulmonary embolism 449
 in syncope 195-9
 Hypothyroidism (see *Myxædema*) 476
 Hypoxia
 electrocardiogram 101
 in pulmonary heart disease 464
 test for angina pectoris 378
 Hysteria 199 263 536
 Iced water 138
 immersion in 528
 Icterus 175
 Idiopathic
 dilatation of the pulmonary artery 21
 hypertrophy of the heart 205
 pulmonary hypertension (see *Primary pulmonary hypertension*) 461
 Idioventricular pace maker 1-3
 Incidence of heart disease 1
 Indifferent electrode 68
 Infection and the heart 313 18
 Infective endocarditis (see *Bacterial*) 350
 Inferior vena cava
 ligation 455
 pressure on 196
 \ ray 25 7 345
 Influenza 313
 Inframammary pain 537
 Infundibular stenosis 229 236
 Initialentricular deflection 67-70
 Injuries to the heart
 direct 525-7
 indirect 528-31
 Innocent
 pain in the chest 537
 parasternal murmur 222
 systolic murmur 281
 Innominate aneurysm 423
 Insomnia 179
 Inspection
 general 2
 of the heart 6
 Interatrial block 84
 Interatrial septum 213
 Intermittent claudication
 in beri beri 516
 in coarctation 208
 pain in 374
 Interpolated extrasystoles 131
 Interstitial myocarditis 315 318
 Interventricular septum
 defect of 220
 displaced 425
 perforated 404
 Intestinal distension 111

- Intracardiac leads 73
 Intracardiac thrombosis 444
 Intra myocardial pressure 433
 Intrathoracic pressure 16
 in emphysema 42
 in left ventricular failure 160
 Intravascular clotting 445
 Intraventricular block 70
 Intrinsic deflection 68
 Intravenous infusions
 in anemia 503
 in circulatory failure 311 314
 and venous pressure 155
 Iodide in syphilitic aortitis 367
 Iodine
 in colloid 48
 lack of 49
 radioactive 191 490
 test for thyrotoxicosis 489
 and thiouracil 492
 in thyrotoxicosis 491
 Iodothyrim 478
 Irritable heart 535
 Ischaemic heart disease (see *Angina pectoris*
 and *Myocardial infarction*) 370-411
 acute coronary insufficiency 384 408 411
 age incidence 371
 alcohol in 371
 angina pectoris 374-86
 definition 370
 etiology 372
 history 370
 incidence 370
 myocardial infarction 386 408
 occupation and 371
 pathogenesis 372-4
 sex incidence 370
 tobacco and 371
 Isolated myocarditis 318-22
 Isometric contraction 19
 Isopotential level 79
 Isopropyl nor adrenalin
 in asthma 473
 in heart block 124
 Isthmus aortic 207
 J and c 175 306
 Joffroy's sign 485
 Jugular phlebogram (see *Jugular pulse*) 19
 Jugular pulsation (see *Jugular pulse*) 169-
 170
 Juxta medullary hypoxia 420 421
 Kahn test 358
 Katharticon 19
 Keith and Flack nodules 108
 Kempner diet 438
 Kent bundle of 141
 Kellin 382
 Kidney (see *Nephritis* and *Renal*)
- Kinked aorta 421 474
 Kinked carotid 423
 Koilonychia in thyrotoxicosis 488
 Kusmaul's sign 349
 Kymogram in cardiac infarction 397
 Kymography 29
 Kyphoscoliosis 474
 Laryngeal palsy
 in aneurysm 361
 in mitral stenosis 293
 Late systolic murmur 282
 Leading questions 1
 Leads (electrocardiographic)
 bipolar 68
 C1 74 77
 chest 68
 C1 74 77
 Goldberger's augmented 75
 intracardiac 73
 oesophageal 73
 standard 77
 unipolar 68
 limb 74
 V 68 70
 Left anterior oblique position 27
 Left auricle (see *Auricle left*)
 Left auricular appendage 26 216
 enlargement 47-52 288
 P wave 83
 pressure 213
 Left axillary node 85 86
 Left bronchus 27
 in aneurysm 360
 in mitral stenosis 293
 Left bundle branch block (see *Bundle
 branch block*) 90 126
 Left inframammary pain 537
 Left ventricle
 abnormalities 42
 in atheroma 34
 X ray 25-7 35
 Left ventricular aneurysm (see *Cardiac
 aneurysm*) 404
 Left ventricular chord 34
 Left ventricular enlargement (in) 42
 aortic incompetence 298
 aortic stenosis 301
 coronary 208
 electrocardiogram 86 87
 familial cardiomegaly 205
 hypertension 423 429
 mitral incompetence 283
 myocardial infarction 377
 patent ductus 224
 tricuspid atresia 230
 ventricular septal defect 222
 X ray 42
 Left ventricular failure 157 (3
 in acute nephritis 327

- Left ventricular failure (contd.)
 aminophylline 190
 in anæmia 503
 in aortic incompetence 229 365
 basal rates 160
 blood pressure 160
 cardiac asthma 160
 cardiac output 160
 Cheyne Stokes breathing 167
 circulation time 161 163
 clinical features 158
 cyanosis in 177
 definition 154-6 157
 digitalis 179-84
 dyspnoea 158
 electrocardiogram 84
 etiology 156 157
 gallop rhythm 164
 hydrothorax 161
 in hypertension 425 435
 intrapleural pressure 160
 low sodium diet 185-9
 lung volume 161
 mechanism 154 158
 mercurial diuretics 184
 morphine 190
 in myocardial infarction 389 403
 orthopnoea 158
 oxygen 190
 P wave changes 84
 paroxysmal cardiac dyspnoea in 158
 posture 158 179
 prognosis 178-9
 pulmonary congestion 158-61
 pulmonary oedema 158-61
 pulsus alternans 166
 in rheumatic carditis 279
 tetraethylammonium bromide 190
 theophylline 190
 treatment 179-91
 venesection 179
 venous pressure 160
 venous tourniquets 190
 vital capacity 158
 X-ray 44 161
- Left atricular preponderance (see *Left atricular enlargement*) 87 426
- Eggs
 blood pressure in 5 208
- Lesser systolic click 165
- Leucocytosis (in) 22
 bacterial endocarditis 322
 myocardial infarction 389
 pulmonary infarction 453
 rheumatic carditis 259
- Lid lag 481 485
- Lid retraction 481 485
- Limb leads 74 77
- Lipoids in atherosclerosis 373
- Liver (see *Hepatic*)
 dipping on 172
 engorgement of 171-2
 palpation of 171
 percussion of 171
 sugar icing of 353
- Long diameter 34
- Lordosis syncope in 196
- Low blood pressure (see *Hypotension*)
- Low output failure 159
- Low sodium diet (in) 185-9
 heart failure 185
 myocardial infarction 406
 hypertension 437
- Low voltage electrocardiogram (in) 103
 anæmia 506
 myxoedema 496
 pericardial effusion 342
 Pick's disease 349
 pulmonary heart disease 467
- Luetic aortitis (see *Syphilitic*) 358-68
- Lugol's iodine in thyrotoxicosis 489 491
- Lumbo dorsal sympathectomy 195 439
- Luminal (see *Phenobarbitone*)
- Lungs (see *Pulmonary*)
 collapse of (see *Collapse*) 35
 cystic 464
 examination of 11
 rheumatic 265
 silicosis of 464
- Lung volume 17-19
 in emphysema 19 472
 in left ventricular failure 19 161
 in mitral stenosis 292
- Lupus disseminatus 315
- Lutembacher's syndrome 216 218 220
- Lymphadenoid goitre 486
- Lymphatic oedema 173
- Machinery murmur (see *Continuous murmur*) 223
- Mackenzie's polygraph 19
- Macula star figure 423
- Magnesium sulphate 139
- Maladie de Roger 222
- Malar flush 286
- Malignant endocarditis 330
- Malignant exophthalmus 481
- Malignant goitre 486
- Malignant hypertension 433
 arteriolar necrosis in 419
 experimental 419
 papilloedema in 424
 prognosis 435
 renal function 433
 retinopathy in 423-4
- Malignant pericarditis 354
- Malignant pulmonary emboli 458
- Malingering 540
- Malnutrition 321

- Manifest mean vectors 82
 Marey's law 110
 Marginate erythema 263
 Massive pulmonary embolism (see *Pulmonary embolism*)
 clinical features 447-9
 definition 447
 electrocardiogram 447-51
 mortality 444 453
 syncope in 194
 treatment 455
 Measurements cardiac 34
 Mecholol (see *Cholinergic drugs*) 138 200 516
 Medial calcification 419 422
 Medial necrosis of the aorta 521
 Mediastinal tumour 34 362
 Mediastinopericarditis 351
 Medico legal aspects 532
 Meniere's syndrome
 auricular fibrillation in 531
 syncope in 200
 Meningitis and auricular fibrillation 143 531
 Meningococcal myocarditis 314 315
 Menopause and hypertension 416 437
 and syncope 200
 and thyrotoxicosis 480
 Mercurial diuretics 184
 Mercuriophylline 184
 Mercury in syphilis 367
 Mersalyl 184
 Methylthiouracil (see *Thiouacil*) 492
 Metre of the heart beat
 amphibrachic 164
 anapestic 165
 dactylic 164
 Metaclavicular line 6
 Mid diastolic murmur 118
 Mitral wheel murmur (see *Continuous murmur*)
 Milroy's oedema 173
 Mirror image dextrocardia 204
 Mitral calcification 63 284 517
 Mitral diastolic murmur 288
 in anaemia 503
 in aortic incompetence (Austin Flint) 296
 in atrial septal defect 216
 in coarctation of the aorta 208
 functional 288
 in mitral stenosis 286-8
 in patent ductus 224
 in rheumatic carditis 270
 in thyrotoxicosis 487
 in ventricular septal defect 222
 Mitral incompetence 290
 and aneurysmal left auricle 284
 clinical features 282
 functional 282
 kymography 284
 left auricle 283
 left ventricular enlargement 283 284
 physiology 282
 rheumatic 282
 traumatic 530
 X-ray 54 283
 Mitral I wave (see *P mitral*) 84 290
 Mitral presystolic murmur 270 288 296
 Mitral ring 282
 Mitral stenosis 284-94
 in active rheumatic carditis 274
 angina pectoris in 274
 angiocardiogram 47 270
 arterial oxygen saturation 272
 and atrial septal defect (Lutembacher's syndrome) 216
 auricular fibrillation in 247 293
 auricular flutter in 293
 bacterial endocarditis in 294
 bronchitis in 293
 calcified 284
 cardiac output 292
 circulation time 292
 clinical features 285
 and coarctation of the aorta 204
 collapse of the lung in 293
 complications 293
 congestive failure in 285 293
 course 294
 cyanosis 285
 development of 258
 dyspnoea 285
 electrocardiogram 290-2
 emboli (systemic) 293
 function studies 292
 haemoptysis 293
 hypertension in 294
 intracardiac thrombosis in 444
 laryngeal pulsus 293
 left auricle in 47 285 288
 left auricular pressure 285
 lung volume 285 292
 mitral diastolic murmur 286-8
 mitral incompetence 284
 opening snap 286
 orthopnoea 158
 P wave 84 290
 pulmonary cardiac dyspnoea 28 293
 physiology 284
 physiology 284
 presystolic murmur 288
 pulsus 294
 pulmonary congestion 285
 pulmonary embolism in 443
 pulmonary haemorrhages 290
 pulmonary hypertension 85 292
 pulmonary incompetence 288
 pulmonary infarction 293

- Mitral stenosis (contd.)
 pulmonary oedema 293
 pulmonary tuberculosis in 294
 rheumatic history 276
 and rheumatoid arthritis 256 294
 surgical treatment 308
 and thyrotoxicosis 294 494
 traumatic 530
 treatment 308
 triple rhythm 166 286
 vaso constriction in 285
 venous pressure 285 292
 vital capacity 285 292
 X ray 56 288
- Mitral systolic murmur 273 282
- Mitral valve disease 280
- Mitral valvulitis 270 273
- Mitral valvulotomy 308
- Möbius sign 485
- Monckeberg's sclerosis 419 422
- Mongolism 93
- Monocardiogram (of Mann) 125
- Morbus cœnuleus 203
- Morphine (in)
 asthma 473
 congestive failure 179
 dissecting aneurysm 523
 myocardial infarction 406
 paroxysmal cardiac dyspnoea 190
- Mortality from heart disease 1
- Müller's experiment 224
- Mural thrombosis (in) 444
 bacterial endocarditis 330 336
 isolated myocarditis 318 321
 mitral stenosis 293 444
 myocardial infarction 388 404
- Murmur (and see under items listed)
 aortic diastolic 295
 aortic systolic 281 301
 apical systolic 273 281-2
 arterio venous (see Continuous) 514
 Austin Flint 296
 basal systolic 281 301-2
 cerebral or cranial 511
 continuous 223 229
 Durozier's 297
 extracardiac 281
 functional 281-2
 Gibson's (continuous) 223
 Graham Steell (pulmonary diastolic)
 216 288
 hæmic 503
 humming top (continuous) 223
 innocent 273 282
 machinery (continuous) 223 229
 Mill wheel (continuous) 223
 mitral diastolic 288
 mitral presystolic 118 270 288 296
 mitral systolic 273 282
 propagation of 9
- Murmur (contd.)
 pulmonary diastolic 216 288
 pulmonary systolic 216 229 231
 respiratory 281
 thyroid 486 493
 to and fro 364
 train in tunnel (continuous) 223
 tricuspid diastolic 306
 tricuspid systolic 304
 vascular 208
 venous humming 229
 whining 331
- Myasthenia gravis 487
- Mycotic aneurysm 333
- Myocardial abscess 354
- Myocardial bruise 529
- Myocardial fibrosis (see Fibrosis of the heart) 307
- Myocardial infarction 386-408
 after effects 405
 and angina pectoris 405
 anterior 389
 antero lateral 391
 antero septal 391
 auricular fibrillation in 389 403 407
 auricular flutter in 389 403 407
 blood pressure 389 408
 bundle branch block in 392
 cardiac aneurysm 388 397 404
 cardiac enlargement 397
 cerebral embolism 404
 cerebral thrombosis 404 406
 clinical features 388-9
 clotting time 443
 complications 397
 congestive failure 403 405
 and coronary thrombosis 386 387 388
 course 397
 diagnosis 405
 dicoumarol 406
 diet 406
 ectopic beats 389 397
 electrocardiogram 93 389-97
 electrokymography 397
 emboli (systemic) 404
 erythrocyte sedimentation rate 38
 etiology 368
 fever 389
 gallop rhythm 389
 heart block in 389 403
 heart sounds 389
 heparin 406
 impending 384
 kymography 44 397
 left ventricular failure 403
 leucocytosis 389
 location of 387
 morphine 406
 mural thrombosis 388 404 445
 myomalacia cordis 398

- Myocardial infarction (contd.)
 nodal rhythm 403
 paroxysmal cardiac dysrhythmia 389
 paroxysmal tachycardia in 389 403
 pathology 389
 perforated septum 404
 perforation of 355
 pericarditis 405
 phlebotrombosis 403
 posterior 94 391
 prognosis 405
 pulmonary embolism 403 405
 quinidine 406
 ruptured heart 353 388 404 405
 signs 389
 site of 387
 symptoms 389
 syncope in 389
 traumatic 531
 treatment 406
 venous pressure 389
 ventricular aneurysm 388 397 404
 ventricular fibrillation 397 403
 X-ray 44 397
- Myocardial ischaemia (see *Ischaemic heart disease*)
- Myocardial necrosis
 from acetylcholine 322
 in acute coronary insufficiency 410
 from digitalis 322
 in diphtheria 312
 ischaemic 388
 in isolated myocarditis 318
 toxic 314
 vagal 322
- Myocarditis 318-28
 allergic 315
 in bacterial endocarditis 315 331
 335
 Chagas disease 315
 clinical features 316-17
 in digitalis poisoning 322 5
 differential 311
 in disseminated lupus 315
 from drugs 326-7
 electrocardiogram in 104 317
 from emetine 325
 Fiedler's 318
 influenzal 316
 isolated 318-21
 mononucleococcal 315
 nephritic 327
 in pneumonia 315
 prognosis 317
 protozoal 315
 pyogenic 322
 rheumatic 266
 in scrub typhus 315
 sulphamides 315
 toxic 314
- Myocarditis (contd.)
 treatment 317
 in typhoid fever 316
- Myocardium nutrition 11
- Myomalacia corli 358
- Myxoedema 416-9
 anaemia in 416
 angina pectoris in 417
 artificial 191
 basal metabolic rate 416
 blood cholesterol 416
 cardiac enlargement 417
 circulation time 416
 clinical features 416
 electrocardiogram 416
 hydropicardium in 356 417
 local (in thyrotoxicosis) 488
 Raynaud's phenomenon 416
 from thiouracil 190 385 412
 treatment 419
- Myxoma auricular 193
- NAB 367
- Natelle & digitaline 148
- Nephritis
 acute diffuse 327
 in bacterial endocarditis 334
 hypertension in 417
 myocarditis in 327
 pulmonary oedema 327
 retinopathy 424
 in rheumatic fever 332
 venous pressure 327
 chronic 433
 cardiac enlargement 415
 hypertension 416 417 418
 oedema 173
 prognosis 435
 retinopathy 423 4 4
 embolic 334
- Neptal 184
- Nerve supply of the heart 109
- Neurolept therapy asthenia (see *Cardiac neurosis*) 535-45
 nomenclature 535
- Neutro electrode 68
- Neutral segment 26
- Nicotine 320
- Nikethamide (coamine) 190 455
- Nitrites 381
- Nitroglycerine 378 381
- Nodal ectopics 128
- Nodal escape 113
- Nodal rhythm 113 117
- Nodules of the heart 108
- Nodular goitre 479 485
- Nodule
 peripheral 257
 rheumatic subcutaneous 265
 rhumatoid 256

- Nor adrenaline in asthma 473
 in heart block 124
 Notching of the R wave 79 90 93
 Notching of the ribs 208
 Novarsurol 184
 Novocaine (intravenous)
 in paroxysmal tachycardia 139
 in ventricular fibrillation 151
 Novurit 184
 Nutritional cardiopathy 321
 in beri beri 314-16
 in hypertension 157
 Ob sity
 in angina pectoris 381
 electrocardiogram 86
 in heart failure 179
 in hypertension 416 436 437
 X ray of heart in 35
 Oblique positions (X ray) 26-7
 Occupation (in angina pectoris) 371
 Ocular fundi (see *Fundi*) 5 423
 Œdema 172-3
 allergic 173
 in anaemia 173 503
 angioneurotic 173 338
 cardiac 173
 cerebral 434
 infra orbital 173
 lymphatic 173
 Milroy's 173
 nephritic 173
 nephrotic 173
 physiology 172-3
 in Pick's disease 349
 pulmonary 158
 renal blood flow in 173
 rheumatic 257
 sodium retention in 173
 starvation 173
 in superior vena cava obstruction 173
 360
 treatment 184 185
 in venous thrombosis 173 453
 Œsophageal lead 73 391
 Œsophageal pain 406
 Œsophageal spasm 377
 Œsophagus
 displacement from
 left auricular enlargement 47 288
 left ventricular enlargement 47
 pericardial effusion 61
 right aortic arch 42 242
 unfolding of the aorta 429
 obstruction of 361
 X ray of 26
 Œstrogens 200
 Oliguria 172 174
 Opening snap 286
 Ophthalmoscopy 5 423
 Opposing movement point of 26
 Orthodiagraphy 27
 Orthopnoea 158
 Orthostatic hypotension 200 538
 Orthostatic syncope 195
 Ortner's sign 293
 Oscillograph 27
 Osler's nodes 333
 Osmotic pressure (in œdema) 173
 Ouabain 148 184
 Overloading of the heart 154
 Oxygen
 arterio venous difference 15
 capacity 16 17
 carrying power of blood 17
 consumption 15 488
 content 16 17
 in cyanosed congenitals 243
 lack 167 201 378
 in pulmonary embolism 455
 in pulmonary heart disease 464 473
 in pulmonary œdema 190
 saturation 16 17
 unsaturation 16
 P inverted 83
 in auricular ectopics 127
 in coronary sinus rhythm 116 136
 in dextrocardia 204
 in nodal rhythm 117
 in paroxysmal auricular tachycardia 134
 135
 I mitrale 84
 in aortic valve disease 299
 in hypertensive heart disease 84
 in left ventricular failure 84
 in mitral stenosis 290
 P pulmonale 83
 in Fallot's tetralogy 242
 and giant a waves 170
 in mitral stenosis 290
 physiology 467
 in pulmonary embolism 450
 in pulmonary heart disease 463 467
 in pulmonary stenosis 231
 in thyrotoxicosis 487 490
 in tricuspid stenosis 307
 P wave
 abnormal 83
 normal 78 467
 physiology 67
 P R interval
 and heart sounds 10
 in myocarditis 312 317 321
 normal 78
 prolonged 118 271 316
 shortened 139
 in rheumatic carditis 271
 Pacemaker 108
 wandering 117

- Jagged disease 517
 Pain
 in angina pectoris 375 376
 in dissecting aneurysm 522
 left inframammary 537
 in massive pulmonary embolism 447
 in myocardial infarction 388
 oesophageal 377
 pericardial 341
 in pulmonary infarction 453
 spinal 377
 Palate high arched 216
 Pallor
 in anaemia 506
 in anxiety states 506
 in hypertension 422
 in myocardial infarction 389
 in paroxysmal cardiac dyspnoea 160
 peripheral 506
 in pulmonary embolism 447
 in vaso-motor syncope 197
 Palmar flush 517
 Palpation (of the heart) 6
 Palpitations (in) 536
 anaemia 503
 anxiety states 536
 auricular fibrillation 147
 ectopic beats 130
 hypertension 422
 paroxysmal tachycardia 131
 thyrotoxicosis 487
 Pancarditis 255
 Pancreatitis 406
 Papilloedema (in)
 acute nephritis 424
 bacterial endocarditis 334
 cerebral tumour 44
 hypertension 424
 pulmonary heart disease 465
 toxæmia of pregnancy 424
 Paracæsthesia pericardial 348
 Paradoxical embolism 457
 in cardiac catheterisation 16 242
 in Fallot's tetralogy 243
 Paradoxical pulsation
 of left auricle 52 283
 of left ventricle 44 397
 Paradoxical pulse 349
 Parasternal distension 360
 Parasternal murmur 222
 Paravertebral injections 386
 Pardee's sign 97 391
 Paroxysmal auricular fibrillation 147
 in myocardial infarction 403
 in thyrotoxicosis 487 493
 Paroxysmal auricular flutter 142
 Paroxysmal bundle branch block 126
 Paroxysmal cardiac dyspnoea (see *Left ventricular failure*) 158-60
 in mitral stenosis 285 291 293
 Paroxysmal heart block 120
 Paroxysmal hypertension 417 421
 Paroxysmal tachycardia 131-41
 auricular 132
 angina pectoris in 132
 classification 131
 clinical features 132-3
 congestive failure 132
 diagnosis 132-3
 in digitalis poisoning 322
 electrocardiogram 132-7
 etiology 132 137
 in heart block 124
 mechanism 133
 in myocardial infarction 403
 in myocarditis 312 317 321
 nodal 132 136
 physiology 132
 prognosis 132 138
 in rheumatic carditis 273
 in Stokes Adams syndrome 123
 supraventricular 132
 syncope in 132 194
 treatment 138
 ventricular 137
 in Wolff Parkinson White syndrome 139 141
 Parry's disease (see *Thyrotoxicosis*) 478
 Partial heart block (see *Heart block*) 118-120
 Paste electrode 68
 Patent ductus arteriosus 223-9
 angiocardigram 224
 bacterial endocarditis in 227
 cardiac catheterisation 224 227
 clinical features 223
 compression of the aorta 209
 differential diagnosis 229
 electrocardiogram 224 226
 embryology 206
 in Fallot's tetralogy 236 243
 hæmodynamics 223
 incidence 223
 prognosis 227
 surgical treatment 228
 surgery 224
 Patent foramen ovale 213
 in Fallot's tetralogy 242
 and paradoxical emboli 457
 in pulmonary hypertension 245 464
 in pulmonary stenosis 231 232
 Pedunculated myxoma 193
 Penicillin
 in bacterial endocarditis 337
 in pyogenic pericarditis 354
 in rheumatic fever 275
 in syphilis 367
 toxic reactions of 338
 Pericarditis of the heart 7
 in urinary 360

- Percussion of the heart (contd.)
 in emphysema 465
 in pericardial effusion 345
- Percussion wave (of the pulse) 20
 in aortic incompetence 96
 in aortic stenosis 300
- Perforation
 of interventricular septum 404
 of myocardial infarction 404
 of sinus of valsalva 523
- Peri arteritis 315 461
- Pericardial adhesions 308 351
- Pericardial calcification 308 349 352
- Pericardial click (or knock) 163
- Pericardial ecchymose 355
- Pericardial effusion 344-8
 cardiac compression (tamponade) 345
 clinical features 344-5
 collapse of lung 345
 differential diagnosis 321 348
 electrocardiogram 99 342-4
 etiology 341
 friction 345
 hæmorrhagic (see *Hæmopericardium*)
 353 354
 in malignant hypertension 356
 in myxœdema 356 494
 paracentesis 348
 percussion 345
 physiology 344-5
 purulent 354
 Q Tc 348 352
 treatment 348
 X ray 61 345
- Pericardial foreign body 355 325
- Pericardial hæmatoma 527
- Pericardial injury 354 525 529
- Pericardiectomy 351
- Pericarditis 341-56
 benign 354
 chronic constrictive (see *Pick's disease*)
 348-51
 dry 341
 electrocardiogram 99 342-4
 etiology 341
 fibrinous 341
 friction 341 345
 malignant 354
 in myocardial infarction 355
 pain 341
 pneumococcal 354
 in polyserositis 353
 pyogenic 354
 recurrent 525
 rheumatic 266 352
 staphylococcal 354
 streptococcal 354
 traumatic 354
 tuberculous 353
 uræmic 355
- Peripheral blood flow
 in arteriovenous aneurysm 514
 in thyrotoxicosis 489
- Peripheral circulatory failure (see *Circulatory failure*) 195 311
- Peripheral cyanosis 177
- Peripheral neuritis
 in beri beri 516
 in diphtheria 311
- Peripheral resistance collapse of 195
- Periodic breathing 167
- Pernicious anæmia (see *Anæmia*) 502
- Persistent truncus arteriosus 250
- Petechiæ (in)
 anæmia 332
 asphyxia 531
 bacterial endocarditis 332
 fat embolism 458
 rheumatic 257 263
 salicylate poisoning 274
 vitamin deficiency 332
- Pethidine 473
- Phenobarbitone 130 437 421
- Phæochromocytoma 416 417 421
- Phlebitis 445 446 453
- Phlebogram 19
- Phlebothrombosis 446 453
 treatment 454
- Phonocardiogram (of)
 aortic incompetence 295
 functional mitral diastolic murmur
 287
 innocent systolic murmur 281
 late mitral systolic murmur 283
 mitral diastolic murmur 286
 mitral incompetence (organic) 293
 opening snap 287
 presystolic murmur 287
 third heart sound 21 164
- Phonocardiography 20-1
- Phosphatase 517
- Photo electric cell 31
- Physiological bundle branch block 140
- Physiological heart block 133 143
- Physiology (see *stem required*)
- Pick's disease 348-51
 cardiac output 347
 clinical features 349 352
 electrocardiogram 99 349
 etiology 349
 pathology 348
 physiology 349
 surgical treatment 351
 X ray 61 349
- Pigeon chest 216
- Pigmentation
 in thyrotoxicosis 488
 in tricuspid stenosis 306
- Pistol shot sound 296
- Pitkin's menstruum 454

- Pitressin test (m)
 berberis 516
 epilepsy 199
 Pituitary irradiation (in thyrotoxicosis) 433
 Plasma infusions 314
 Plasma loss of 195
 Plasma proteins (in edema) 173
 Plateau pulse 300
 Pleural venous drainage of 161 174
 Pleural effusion (see *Hydrothorax*)
 haemorrhagic 453
 rheumatic 265
 venous pressure in 171
 Pleurisy
 in pulmonary infarction 447 453
 rheumatic 265
 Pleuro-pericardial adhesions 164
 Pneumomediastinum 464
 Pneumonia
 myocarditis in 314 315
 pericarditis in 354
 rheumatic 265
 rhythm changes in 143
 Pneumothorax systolic click 165
 Point of opposing movement 26 34 42
 Polarity of electrocardiogram 68
 Polarised cell 65
 Polyarthritides
 allergic 256
 dysenteric 256 262
 gonococcal 256
 rheumatic 265
 rheumatoid 256
 Polychromasia (in heart failure) 22 175
 Polycythemia (in)
 arteriovenous neurysm of the lung 513
 Ayras disease 474
 congenital heart disease 22 177
 congestive heart failure 22
 Fallot's tetralogy 237
 pulmonary heart disease 22 462 464
 Polygraph 19
 Polyserositis 353
 Polyuria 433
 Postural hypotension 195 200
 Postural syncope 136
 Potent effect on
 angina pectoris 376
 apex beat 351
 blood pressure 5
 cardiac output 16
 electrocardiogram 86 87
 left ventricular failure 159
 rheumatic carditis 275
 right auricle pressure 158 179
 venous pressure 169
 vital capacity 158
 Potassium 326
 in ectopic beats 130
 electrocardiogram 105 326
 Ictadium (contd.)
 poisoning 326
 in uraemia 105
 and ventricular fibrillation 127
 Ictadium iodide in syphilis 317
 Ictid operation 243
 Precordial defibrity 6
 Ictid leads 68
 Ictid excitation (W I W syndrome) 133 41
 Ictid pregnancy 507-11
 bacterial endocarditis in 510
 congenital heart disease in 508
 electrocardiogram 508
 hypertension in 511
 physiology 307
 rheumatic heart disease in 508
 syncope in 196
 and thyrotoxicosis 495 510
 Ictid beats (see *Ectopic beats*) 127
 Ictid agent 418 0
 Presystolic gallop (see *Cellular rhythm*)
 164
 Presystolic murmur
 Austin Flint 296
 in mitral stenosis 288
 Ictid pulmonary hypertension 461-4
 clinical features 462
 diagnosis 463
 electrocardiogram 463
 etiology 461
 incidence 462
 pathology 461
 prognosis 464
 physiology 462
 treatment 464
 X rays 57 463
 Primary vascular sclerosis 461
 Pocane penicillin 338
 Propylthiouracil (see *Thiouacil*) 19 492
 Prothrombin 138
 Protrombin time 454
 Protodiastolic gallop (see *Gallop rhythm*)
 156
 Protozoal myocarditis 315
 Pseudoerythroidism of the liver 348
 Psychogenic faint 196 538
 Psychomotoros (see *Cardiac neurosis*) 535
 545
 Psychosomatic signs and symptom 536-
 540
 Pulmonary apoplexy 293 513
 Pulmonary artery 26
 Pulmonary retrovascular aneurysm 513
 Pulmonary artery
 abnormalities of 52
 neurysm of 475
 angiosarcoma (in rim) 31
 atherosclerosis 461
 chronic thrombosis 15
 development 206

Pulmonary artery (contd.)

- dilatation (in) 52
 - atrial septal defect 52 18
 - beri beri 57 515
 - Eisenmenger's complex 52 244
 - idiopathic 212 475
 - mitral stenosis 56 288
 - normal variation 231
 - patent ductus 52 224
 - primary pulmonary hypertension 57 463
 - pulmonary heart disease 57 472 475
 - pulmonary stenosis 52 231
 - syphilis 475
 - transposition 246
 - ventricular septal defect 56 22
 - X ray 52 7
- hypoplasia of 57 240
- palpation of 7
- pressure (in)
 - atrial septal defect 18
 - Bernheim's syndrome 425
 - Eisenmenger's complex 245
 - emphysema 472
 - Fallot's tetralogy 242
 - hypertension (essential) 425
 - idiopathic dilatation of the pulmonary artery 212
 - mitral stenosis 285 292
 - normal 16
 - patent ductus 227
 - physiology 462
 - Sick's disease 349
 - primary pulmonary hypertension 463
 - pulmonary heart disease (anoxic) 472
 - pulmonary stenosis 231 232
 - ventricular septal defect 220
- pulsation 6
- thrombosis of 447 475
- X ray (normal) 25-7 35
- Pulmonary atresia 204 236 50
- Pulmonary blood flow
 - diminished in
 - Eisenmenger's complex 245
 - Fallot's tetralogy 236 240
 - pulmonary atresia 250
 - pulmonary stenosis with reversal in tetralogy shunt 232
 - tricuspid atresia 250
 - increased in
 - atrial septal defect 216 218 220
 - patent ductus 223 224 227
 - perforated aortic sinus into right side of heart 524
 - transposition 246 250
 - ventricular septal defect 221 222
- Pulmonary blood pressure (see *Pulmonary artery pressure*)
- Pulmonary circulation
 - physiology 446-7 461-2 464

Pulmonary circulation (contd.)

- time (see *Circulation time*) 12-13 161
- Pulmonary congestion
 - in hypertension 425
 - in left ventricular failure 158-61
 - in mitral stenosis 53 288
 - X ray 53 161
- Pulmonary diastolic murmur (see *Pulmonary incompetence*)
- Pulmonary embolectomy 455
- Pulmonary embolism 444-9
 - air 457
 - in bacterial endocarditis 333 445
 - blood pressure in 448 449
 - classification 444
 - clinical features 447 9
 - electrocardiogram 449-51
 - etiology 444-6
 - fat 457
 - foreign body 458
 - hemodynamics 446-7
 - incidence 444
 - in isolated myocarditis 321 445
 - malignant 458
 - massive 194 447
 - in mitral stenosis 293 444
 - in myocardial infarction 403 4 406-7 445
 - pain in 447 453
 - pathology 446
 - and phlebothrombosis 445-6 453
 - prognosis 453
 - and pulmonary infarction 447 453
 - recurrent 449 461
 - septic 457
 - sudden death, 449
 - syncope 194 449
 - in thyrotoxicosis 493
 - and thrombo-phlebitis 445
 - treatment 454-7
 - venous pressure 448 449
 - X ray evidence of 453
- Pulmonary hemosiderosis 290
- Pulmonary heart disease
 - anoxic 464-75
 - aneurysm of the pulmonary artery in 475
 - arterial oxygen 464 472
 - and Ayerza 474
 - carbon dioxide retention 464
 - cardiac output 156 464 465 472
 - clinical features 464-5
 - congestive failure in 156 465
 - cyanosis in 464 465 472
 - definition 464
 - diagnosis 472
 - electrocardiogram 83 465-75
 - emphysema in 465
 - etiology 464
 - incidence 464

- Pulmonary heart disease
 anoxic (cont'd)
 in kyphoscoliosis 474
 papilloedema in 465
 pathology 464
 polycythæmia 465
 prognosis 473
 pulmonary artery pressure 464 472
 residual air 472
 treatment 473
 vaso-dilatation 465
 vaso-motor collapse 465
 venous pressure 465 472
 vital capacity 464 472
 X ray 57 472
 hypertensive (see *Primary pulmonary hypertension*) 461
 subacute 449 459 461
 Pulmonary hypertension 461
 in atrial septal defect 216
 Bilharzial 461
 in Eisenmenger's complex 245
 idiopathic 461 463
 in mitral stenosis 285 292
 in patent ductus 2
 in periarteritis 461
 primary 461 463
 in pulmonary heart disease (anoxic) 42
 in recurrent pulmonary embolism 461
 second heart sound in 10 462
 in ventricular septal defect (cont'd)
 Pulmonary hypoxia 464
 Pulmonary incompetence (functional)
 atrial septal defect 216
 Eisenmenger's complex 244
 idiopathic dilatation of the pulmonary artery 212
 mitral stenosis 288
 patent ductus 223
 pulmonary embolism 449
 pulmonary hypertension 462
 ventricular septal defect 222
 Pulmonary infarction (see *Pulmonary embolism*)
 in bacterial endocarditis 333
 clinical features 453
 in congestive heart failure 174
 hydrothorax in 174
 in isolated myocarditis 321
 jaundice in 175
 in mitral stenosis 293
 pathology 447
 X ray 453
 Pulmonary ischæmia (see *Pulmonary blood flow*)
 Pulmonary oedema (see *Left ventricular failure*) 158-62
 acute nephritis 327
 anæmia 503
 QX
 Pulmonary oedema (c ntd)
 hypertension 425 435
 mitral stenosis 293
 myocardial infarction 389 403
 rheumatic carditis 20
 ruptured aortic cusp 530
 treatment 189-90
 X ray 44 161
 Pulmonary oligæmia (see *Pulmonary blood flow diminished*)
 Pulmonary P wave (see *P pulmonale*) 83 467
 Pulmonary pleonæmia (see *Pulmonary blood flow increased*)
 Pulmonary plethora (see *Pulmonary blood flow increased*)
 Pulmonary second sound (see *Second heart sound*) 10
 Pulmonary stenosis (excluding Fallot's tetralogy) 229-36
 angiocardigram 234
 and atrial septal defect 231 232
 bacterial endocarditis in 231 330
 cardiac catheterisation 232
 classification 203 232
 clinical features 229
 cyanosis in 232
 diagnosis 231 232
 electrocardiogram 231
 infundibular 229 232
 interatrial reversed shunt in 232
 interventricular reversed shunt in 232
 patent foramen ovale in 232
 pathogenesis 229
 prognosis 231
 subvalvular 229 232
 surgical treatment 231 236
 syncope 232
 valvular 229 232
 ventricular septal defect in 231 232
 X ray 52 231 232
 Pulmonary systolic murmur (in)
 atrial septal defect 216
 Fallot's tetralogy 237
 functional 231
 patent ductus 223
 pulmonary stenosis 229
 ventricular septal defect 222
 Pulmonary thrombosis 447 475
 Pulmonary tuberculosis (n)
 Fallot's tetralogy 243
 pulmonary heart disease 464
 pulmonary stenosis 231
 Pulmonary valvulotomy 231 236 244
 Pulmonary vascular sclerosis 461
 Pulmonary vasoconstriction 462
 Pulmonary veins anotomy of 234
 Pulmonary venous congestion (see *Pulmonary congestion*) 160-1 288
 Pulmonary venous samples 213 234

- Pulse (and see *Water hammer*)
 alternation 166
 anacrotic 3 300
 in aortic incompetence 295 296 297
 in aortic stenosis 300
 in atherosclerosis 419
 bigeminal 129 300
 bisferiens 300
 capillary (see *Capillary pulsation*) 297
 in coarctation 3 208
 collapsing (see *Water hammer*) 122 295 297
 Corrigan (see *Water hammer*) 297
 delayed 3 208
 dirotic 314
 double 300
 in hyperkinetic circulatory states 502
 hypertensive 422
 jugular (see *Venous pulse*) 169-71
 paradoxical 349
 plateau 300
 pressure 5 487 50
 rate 110-113
 slow rising, 300
 small (in)
 aortic stenosis 300
 atrial septal defect 216
 circulatory failure 311
 mitral stenosis 286
 myocarditis 312 317 318
 pericardial effusion 345
 Pick's disease 349
 pulmonary embolism 449
 syncope 198
 vaso constriction 416 422
 ventricular septal defect 222
 sustained 300
 velocity 20
 venous (see *Venous pulse*)
 water hammer (see *Water hammer pulse*)
 3 206
 waves 20
 weakness (unilateral) 3
Pulsus alternans 166-7
Pulsus tardus 300
 Pupils (in)
 Cheyne Stokes breathing 168
 Horner's syndrome 361
 epilepsy 198
 syncope 198
 Purkinje fibres 108
 Purpura rheumatica 257
 Pyelonephritis 416 418 436
 Pyogenic myocarditis 322
 Pyogenic pericarditis 354
 Pyridoxin (in thyrotoxicosis) 421
 Pyrogens 17
- Q wave
 in myocardial infarction 97 389
- Q wave (contd)
 normal 78
 in obesity 86
 physiology 69
 in pregnancy 86
 in pulmonary embolism 449 450
- QRS complex
 abnormalitie 85-93
 area 81
 axis deviation 85-87
 basic patterns 73-4
 in bundle branch block 90-3
 in chest leads 70 73
 in left ventricular preponderance 87
 in myocardial infarction 93-4 97 389-397
 normal 70 78
 notched 90
 physiology 67 68-70
 in potassium poisoning 105 326
 in pre excitation 139
 in right ventricular dominance 88
 in ventricular ectopic beats 129
 in ventricular tachycardia 137
 widened 89 105
- QRS vector 82
- QT interval 79
- QTc (in) 79
 digitalis therapy 325
 hypocalcaemia 326
 pericardial effusion 348
 potassium poisoning 105
 rheumatic carditis 71
 uraemia 326
- Quincke's oedema 173
 from penicillin 338
 from quindine 149
- Quindine
 action of 149
 in auricular fibrillation 148
 in auricular flutter 145
 dose 139 149
 in ectopic beats 130
 hypersensitivity 149
 intravenous 139
 in myocardial infarction 151 406
 in paroxysmal tachycardia 139
 risks of 149
 in ventricular fibrillation 151
- R wave (see *QRS complex*)
- Rat bit heart rate in 110
- Radial pulse 3
- Radio active iodine
 for artificial myxoedema 191
 in thyrotoxicosis 490 493
- Radio active sodium 13
- Radiology of the heart (see *X rays* also item or technique required) 25-63
- Radio opaque substances 31

- Radium C 163
 Rales basal 170
 Rapid ventricular filling 156 280
 Rat heart rate in 110
 Rate output curve 178
 Raud's phenomenon (in myxedema) 436
 Read's formula 488
 Recurrent laryngeal palsy
 in aneurysm 261
 in mitral stenosis 293
 Reduplicated heart sounds 164
 Regression wave 67
 Renal blood flow (in congestive failure) 173
 Renal function tests 22
 Renal hypertension 418 419 421 436
 Renal infarction 334
 Renin 419-20
 Reserve air 17
 Residual air 17
 in emphysema 472
 in left ventricular failure 161
 Respiratory
 centre (in Cheyne Stokes breathing) 168
 failure (in pulmonary embolism) 455
 murmurs 281
 sinus arrhythmia 109
 Rest (see *Bed rest*)
 Reticuloecytosis (in congestive failure) 22 175
 Retinopathy
 in bacterial endocarditis 334
 diabetic 424
 hypertensive 423-4
 Retinoscopy 5 423
 Retrograde aortography 209
 Retrograde heart block 117 128
 Rheumatoid arthritis 257 285
 Rheumatoid carditis (and see *Rheumatic fever*) 266-74
 aortic diastolic murmur 270
 auricular fibrillation in 273
 cardiac enlargement 266
 congestive failure 270
 course 275
 diagnosis 273
 electrocardiogram 270
 gallop rhythm in 273
 mitral diastolic murmur 270
 in mitral stenosis 293 294
 mitral systolic murmur 273
 mortality 276
 pathology 57
 pulmonary heart block 27
 pericarditis 266 352
 Q.Tc 71
 relapse 275
 treatment 275
 Rheumatic fever 255-77
 A.C.T.H. in 275
 Rheumatic fever (contd)
 and cardiac neurosis 21
 chorea in 261 275
 clinical features 259
 cortisone in 275
 course 274
 erythema marginatum 263
 erythrocyte sedimentation rate 274
 etiology 255-7
 heparin tolerance 266
 inflammatory features (non specific) 259
 incidence 255
 nodules 265
 pathology 257
 petechiae 263
 pleurisy 265
 pneumonia 265
 polyarthritis 260
 prognosis 276
 recurrence 275
 and rheumatoid arthritis 256
 salicylates 261 274
 skin lesions 263
 streptococcal relationship 255 260
 treatment 274
 Rheumatic heart disease inactive (see under *Latent lesion*)
 adherent pericardium 308
 aortic incompetence 294-9
 aortic stenosis 299-303
 incidence 280
 of valve lesions 280
 mitral incompetence 280-4
 mitral stenosis 284-94
 myocardial fibrosis 307
 rheumatic history in 280
 treatment 308
 tricuspid incompetence 303-6
 tricuspid stenosis 306-7
 Rheumatic state nature of 255
 Rheumatoid arthritis 264
 Rhythm change (see particular rhythm required) 108-51
 Rb notch 208
 Rcdt 185 438
 Rding 236
 Rdl disease 486
 Right anterior oblique position 26
 Right ucle (see *Auditory right*)
 Right auricular pressure (normal) 16 213
 r is d (see *Normal pressure*)
 Right axis deviation 87
 Right bundle branch block (see *Bundle branch block*) 93 26
 split second sound in 126
 Right diastolic 42 236 242
 Right ventricle
 conus 27 47
 undeveloped -49 250

- Right ventricular enlargement (in) 57-9
 atrial septal defect 12 218
 beri beri 515 516
 cardiac impulse in 16 237 462
 cœur en sabot in 57 240
 Ebstein's disease 203
 Eisenmenger's complex 244
 electrocardiogram 88-9 232 463
 Fallot's tetralogy 57 240
 mitral stenosis 285
 pulmonary incompetence 212
 pulmonary heart disease 57 472
 pulmonary hypertension 462 463
 pulmonary stenosis 52 231 232
 right bundle branch block 126 218
 subacute cor pulmonale 459 499
 tricuspid incompetence 303
- Right ventricular failure (in) 168-91
 atrial septal defect 216
 beri beri 515
 massive pulmonary embolism 449
 pulmonary heart disease 462 465
 pulmonary hypertension 462
 pulmonary stenosis 229
 subacute cor pulmonale 449
- Right ventricular pressure
 in Fallot's tetralogy 242
 normal 16
 in pulmonary stenosis 231 232
 raised (see *Pulmonary hypertension*)
- Rigors
 and blood pressure 416
 in cardiac catheterisation 16
 in paroxysmal cardiac dyspnoea 293
 and venous pressure 160
- Ring second sound 359
- Roentgen rays (see *X rays*)
- Roesler's sign 208
- Root pains 361
- Rotation of the heart (and see *Clockwise rotation*) 35
 anti clockwise 73 83 85
 clockwise 70 82 85
- Rotch's sign 345
- RS T segment
 abnormalities of 93-105
 depression of
 in acute coronary insufficiency 410
 in angina pectoris 101 378
 in anoxic states 101 102 410
 from digitalis 101 325
 in left bundle branch block 90
 in left ventricular preponderance 87 426
 elevation of
 in left bundle branch block 90
 in myocardial infarction 94 389
 in pericarditis 90 342
 reciprocal 88
 normal 79
- Rupture of aorta
 in coarctation 211
 in dissecting aneurysm 522
 traumatic 528
- Rupture of aortic aneurysm 360 363
- Ruptured aortic cusp
 in bacterial endocarditis 331
 traumatic 530
- Ruptured aortic sinus 523
- Ruptured chordæ tendineæ 530
- Ruptured heart
 in cardiac aneurysm 404 405
 in myocardial infarction 404
 traumatic 528
- Ruptured mitral valve 530
- Ruptured pulmonary artery 475
- Rutin 332
- S wave (see *QRS*)
 normal 74 79
 physiology 69
- Saccharin circulation time 13 31 161
- Saccular aneurysm (see *Aneurysm*) 38 359 364
- Salicylates 261 274
- Saline
 infusions 503
 manometer 11 12 13
- Salt restriction (see *Sodium*) 185
- Salyrgan 184
 scarlet fever 256
- Scoliosis 6 35
- Scrub typhus 315
- Second heart sound 10 163
 absent
 in aortic stenosis 301
 in emphysema 465
 accentuated aortic element
 hypertension 424
 accentuated pulmonary element
 Eisenmenger's complex 44
 mitral stenosis 286
 patent ductus 224
 pulmonary hypertension 462
 ventricular septal defect 222
- ringing 359
- single
 aortic stenosis 301
 Fallot's tetralogy 240
 pulmonary atresia 250
 pulmonary stenosis 229
- widely split
 atrial septal defect 216 218
 right bundle branch block 126
- Second oblique position 27
- Second ventricular deflection 67 73
- Secondary inversion of T wave 87 89
- Sedatives (in)
 congestive heart failure 172
 ectopic beats 130

- Sedatives (in) (contd)
 hypertension 437 440
 myocardial infarction 406 407
 pulmonary heart disease 473
 thyrotoxicosis 491
 Sedimentation rate (see *Erythrocyte sedimentation rate*) 22
 Semi beats of Stokes 122
 Septal infarction 392
 Septic endocarditis 330
 Septic myocarditis 322
 Septic pericarditis 354
 Septic pulmonary infarction 453
 Septum
 inter auricular 213
 primum 213
 secundum 213
 Serum
 albumin 173
 sickness 311
 Shifting nodal rhythm 117
 Shock 195
 and coronary insufficiency 408
 in dissecting aneurysm 522
 electrocardiogram 410
 in myocardial infarction 389
 in pulmonary embolism 447
 Short P R interval 139
 Short Q T interval 325
 Shrapnel wounds of the heart 525
 Shunt (see *Arteriovenous or Venous arterial*)
 detection 16 511 513
 measurement 221 42
 Sigs 536
 Silicosis 464
 Silver wire artery 423
 Single second sound (see *Second heart sound*) 240
 Sino-auricular block 114-15
 Sino auricular node 108 109 387
 Sinus arrhythmia 109
 Sinus bradycardia 112-13
 in acute hypertension 110
 in athletes 35 110
 cardiac enlargement 35
 in myocardial infarction 389
 in myxoedema 496
 in syncope 195 197
 Sinus node 108
 Sinus rhythm 108
 Sinus tachycardia 110-1
 in anxiety 537
 in congestive heart failure 178
 differential diagnosis 111 133 142
 effect on the heart 12
 in hyperkinetic circulatory states 502
 in myocarditis 312 317 318
 in pericardial effusion 345
 physiology 110-11
 in Packer disease 349
 sinus tachycardia (contd)
 in pulmonary embolism 449
 in rheumatic carditis 253 261
 in thyrotoxicosis 487
 sinus of valsalva perforation of 229 523-5
 situs inversus 204
 skin resistance 68
 sliding nodal rhythm 117
 slow rising pulse 300
 slurring of QRS 90
 Smithwick operation 439
 and post-operative hypotension 125
 smoking (see *Tobacco*)
 sodium (see *Low sodium diet*) 18 -9
 amylal 440
 benzoate 338
 cyanide 12
 dehydrocholate (decholin) 12 161
 retention 173 328
 salicylate 261 274
 soldier's heart (see *Cardiac neurosis*) 535
 sounds (see *Heart sounds*) 10 163
 South American trypanosomiasis 315
 Southey's tubes 189
 specific aortitis (see *Syphilitic*) 358
 specific gravity
 of pleural fluid 174
 of urine 433
 sphygmomanometry 3-5 416
 spider naevi 517
 spinal deformity 474
 spinal ligamentous pain 377
 syphilitic aortitis (see *Syphilitic*) 358
 spirometer 17
 splanchic denervation 439
 spleen enlargement of
 in bacterial endocarditis 332
 in congestive failure 171
 splenic infarct 332
 splinter haemorrhage 332
 split heart sounds (and see *Second heart sound*) 10 163 164
 squinting 237
 ST segment (see *RS T segment*)
 stab wound of the heart 354 527
 standard leads 77
 standardisation of electrocardiograph 68
 staphylococcal
 endocarditis 330
 myocarditis 354
 pericarditis 354
 resistance to penicillin 337
 starling's curve 155
 starling's law 154 295
 starvation oedema 173
 strutius anginosus 366 380 384
 strutius lymphaticus 528
 stellate ganglionotomy 386
 strutius anginosus 485
 sternum depression of 35

- Stethoscopes 7-8
 Stilboestrol 200
 Still's disease 265
 Stokes Adams attacks 122
 Strained heart 542
 Streptococcal
 endocarditis 330
 myocarditis 315
 pericarditis 354
 polyarthritis 360
 Streptococcus and rheumatic fever 255-6
 260-1
 Streptomycin
 in bacterial endocarditis 338
 toxic effects of 338
 in tuberculous pericarditis 353
 Stretch receptors 285
 String galvanometer 67
 Stroke volume (in)
 aortic incompetence 294
 aortic stenosis 299
 atrial septal defect 216
 bradycardia 35
 complete heart block 12
 congestive failure 145
 mitral incompetence 282
 mitral stenosis 285
 patent ductus 223
 pericardial effusion 344
 Pick's disease 349
 premature ectopic beats 129
 tachycardia 110
 thyrotoxicosis 480
 tricuspid stenosis 306
 Strophantidin
 in auricular fibrillation 148
 in heart failure 184
 in pulmonary embolism 455
 Sturge's disease 411
 Subacute bacterial endocarditis (see *Bacterial endocarditis*) 330
 Subacute pulmonary heart disease 449 459
 Subacute rheumatism 260 331
 Subaortic stenosis 212
 Subaortic window 27
 Subarachnoid hæmorrhage 211 434 511
 Subclavian artery
 development of 206
 variations of 211
 Subcutaneous nodules 58 266
 Subintimal hæmorrhage 374 384
 Substernal goitre 485
 Substernal pain 375-7
 Subtotal thyroidectomy 490-1
 Sudden death (in or from)
 angina pectoris 150
 aortic stenosis 150 193 300
 blows 528
 cardiac syncope 193
 chloroform 326
 Sudden death (contd.)
 coarctation of the aorta 212
 digitalis 150 322
 diphtheria 150 312
 dissecting aneurysm 522
 indirect trauma 528
 mercurial diuretics 151 184
 myocardial infarction 397
 myocarditis 315 321
 potassium 326
 pulmonary embolism 449
 rheumatic carditis 277
 ruptured aneurysm 363
 ruptured heart 528
 Stokes Adams syndrome 122 124 193
 syphilitic aortitis 363 365
 trauma 525
 ventricular fibrillation 150 193 397
 Sugar feed liver 353
 Sugar tolerance
 in thyrotoxicosis 488
 Sulphonamide myocarditis 315
 Sulphonamides
 in bacterial endocarditis 336
 in rheumatic fever 277
 Summation gallop 165
 Superior vena cava
 angiocardiogram 34
 obstruction 170 360
 X ray 25-7 59
 Supra aortic triangle 27
 Suprachol (see *Decholin*) 12
 Supra renal
 failure 311
 tumour 421
 Supraventricular tachycardia 132
 Surgical kidney 416 418 436
 Surgical treatment of
 aneurysm 363-4
 angina pectoris 386
 arteriovenous aneurysm 514
 cardiac compression 527
 cardiac failure (by thyroid ablation) 190
 coarctation of the aorta 211
 Fallot's tetralogy 243
 gunshot wound of the heart 526
 hæmopericardium 355 521
 hypotension 439-40
 mitral stenosis 308
 patent ductus 228
 Pick's disease 351
 pulmonary atresia 250
 pulmonary congestion 308
 pulmonary embolism 455
 pulmonary stenosis 231 236
 stab wounds of the heart 524
 thyrotoxicosis 490
 tricuspid atresia 250
 Sweating
 emotional 538

- Sweating (contd.)
 in myocardial infarction 38)
 in pulmonary embolism 449
 thyrotoxic 538
 in vasovagal syncope 198 199
- Sweet clover disease 454
- Sydenham's chorea 262
- Symballophone 8
- Sympathectomy
 in angina pectoris 386
 in hypertension 43)
- Syncope 193-201
 anoxic 201
 in aortic incompetence 194
 in aortic stenosis 194 299
 in auricular fibrillation 147
 in ball thrombus 193
 cardiac 193-4
 in cardiac compression 194
 from cardiac standstill 193
 carotid sinus 196-7
 cerebral 197 200
 from chemical agents 195 196 200
 clinical features 197
 differential diagnosis 197-9
 electrocardiogram 102 410
 etiology 193 194 195 200 201
 from hemorrhage 195
 hyperventilation 200
 in lordosis 472
 in massive pulmonary embolism 449
 194
 mechanism 193 195 200 201
 in myocardial infarction 389
 orthostatic 195
 in paroxysmal tachycardia 132 194
 in pregnancy 196 507
 in pulmonary stenosis 22)
 psychogenic 196 538
 in Stokes Adams syndrome 122
 treatment 200
 vasomotor 101 194 197
 vasovagal 194 197
 from ventricular fibrillation 150 193
- Syphilitic aneurysm 359-64
 abdominal 361
 angiocardigram 363
 of the arch 360
 of the ascending aorta 359
 calcification 363
 clinical features 359-61
 complications 360 363
 course 363
 fusiform 38 359 364
 incidence 359
 prognosis 363
 rupture 363
 saccular 38 359
 treatment 363
 X ray 38 361-3
- Syphilitic angina pectoris 364
 clinical features 364
 course 366
 pathology 366
 physiology 366
 treatment 367
- Syphilitic aortic incompetence 364-5
 clinical features 364-5
 course 365
 electrocardiogram 365
 erythrocyte sedimentation rate 365
 incidence 358 364
 pathology 364
 X ray 364 365
- Syphilitic aortitis (see Syphilitic aneurysm
 angina and aortic incompetence)
- angiocardiology in 363 364
 an a congenital syphilis 358
 confirmatory findings 358
 erythrocyte sedimentation rate in 358
 365
 incidence 358
 Kahn test in 358
 occurrence 358
 pathology 358
 treatment 367-8
 Wasserman reaction in 358
 X ray 39 361-3 364
- Syphilitic endocarditis 358
- Syphilitic heart block 367
- Syphilitic myocarditis 358
- Syphilitic pulmonary arteritis 474
- Systolic blood pressure (see Blood pressure)
 5
- Systolic click 156
- Systolic expansion of the left auricle 52
 283
- Systolic gallop 165
- Systolic indrawing 6
 in adherent pericardium 351 352
- Systolic murmurs (see Aortic Mitral etc.)
 273 281 503
- T wave
 abnormality of 93-105
 accentuated n
 post myocardial infarction 391
 potassium poisoning 104 326
 uræmia 104 3 6
 inverted in
 anemia 506
 aortic stenosis 301
 carbon monoxide poisoning 10 410
 digitalis therapy 101 323
 diphtheritic myocarditis 312
 hypertensive heart disease 426
 myocardial infarction 93 389-97
 myocarditis 104 317 318
 myxœdema 103 494
 nephritis (acute) 327

- T waves
 inverted in (contd)
 pericarditis 99 342-4
 Pick's disease 349
 pulmonary embolism 449-51
 pulmonary hypertension 463
 pulmonary stenosis 231 42
 normal 73-4 79
 physiology 67
 Ta wave 67
 Tachycardia (see *Sinus tachycardia*) 110-112
 Tachypnoea 536
 Tamponade (see *Cardiac compression*) 345 348
 Tawara's node 168
 Teleradiography 29
 Tetany hyperventilation 201
 Tetraethylammonium bromide
 in hypertension 422
 in left ventricular failure 190
 in syncope 195
 Tetralogy of Fallot (see *Fallot's tetralogy*) 236-44
 Theobromine 185
 Theophylline ethylene diamine (see *Amino phylline*) 190
 Thiamine 515
 Thiouracil
 in angina pectoris 384
 in artificial myxoedema 190 385
 in congestive heart failure 130
 in pulmonary heart disease 474
 in thyrotoxicosis 491
 toxic effects 492
 Thiocyanates 438
 Third heart sound
 in mitral stenosis 286
 normal 163 164-5 166
 physiology 165
 in Pick's disease 349
 in pregnancy 307
 in proto diastolic gallop 164
 in summation gallop 165
 Thoracic deformity
 from cardiac enlargement in childhood 6
 depressed sternum 35
 displaced heart 6 35
 kyphoscoliotic 474
 pigeon chest 216
 scoliotic 6 35
 Thrills (see *Related murmurs*)
 detection of 6
 Thromboembolism 444
 Thrombophlebitis 445
 clinical features 453
 treatment 454
 Thrombosis (see *vessel or territory involved*)
 Thyroglobulin 478
 Thyroglossal cyst 487
 Thyroid extract (in myxoedema) 499
 in thyrotoxicosis 493
 Thyroid gland
 enlargement (see *Goitre*) 479
 histology 479
 total ablation 190 384
 Thyroid hormone nature of 478
 Thyroidectomy
 for angina pectoris 384
 for congestive failure 190
 for thyrotoxicosis 490
 Thyroiditis 486
 Thyrotoxic crises 493
 Thyrotoxic heart disease 478
 pathology 480
 Thyrotoxicosis 478-96
 angina pectoris in 495
 auricular fibrillation in 487 493
 basal metabolic rate 488
 blood cholesterol 488
 cardiac enlargement 487
 cardiac output 489
 cardiovascular signs 487
 circulation time 489
 clinical features 480-8
 congestive failure in 487 493
 contributory etiological factors 480
 decalcification of bone 488
 diagnosis 488-9
 electrocardiogram 487 490
 etiology 477
 eye signs 481 5
 goitre 485-7
 hands 485
 historical note 478
 hyperkinetic circulation 487
 and hypertension 494
 iodine in 489 491 492
 local myxoedema in 488
 and mitral stenosis 494
 and myasthenia gravis 487
 pathology 479-80
 peripheral blood flow 489
 physiology 480
 pigmentation 488
 pituitary irradiation for 493
 and pregnancy 495
 prognosis 495
 and pulmonary embolism 493
 radio active iodine in 490 493
 relapse 492
 and rheumatic heart disease 494
 tachycardia 487
 thiouracil in 491-3
 thyroidectomy for 490-3
 and tonsillitis 494
 treatment 490-4
 urinary creatine test 487
 X ray appearances 487
 X ray therapy 493

- Thyrotropic hormones 479 481
 Thyroxine 4, 8 480
 Tics 261
 Tic tac rhythm 178
 Tidal air 17
 Tidal wave 20
 Time-concentration curve 13
 Time marking 67
 Tinnitus 274
 Tobacco 130 326 381
 Todd units 261
 Tonsillectomy
 in rheumatic fever 275
 in thyrotoxicosis 494
 Tonsillitis in rheumatic fever 260
 Tortuosity of the aorta 39
 Toxæmia of pregnancy 327 417
 Toxicomyocarditis (see *Myocarditis*) 311 314
 Tracer substances
 iodine 490
 sodium 13
 Tracheal displacement
 from aneurysm 361
 from goitre 485
 Tracheal obstruction 361
 Tracheal tug 361
 Transition zone 70
 Transposition of the great vessels 246-50
 Transudate 174
 Transverse diameter of the heart 34
 Transverse position of the heart
 electrocardiogram 77
 X ray 35
 Traumatic lesions of the heart 521-32
 angina pectoris 531
 auricular fibrillation 531
 from blast 355
 from blows 355 528 529 530 531
 crush injuries 355
 from direct trauma 5 5-7
 from electric shock 531
 from falls 528 530
 gunshot wounds 354 525-7
 hæmopericardium 354 525 529
 from head injuries 531
 heart block 531
 from indirect trauma 528-532
 medico legal aspects 532
 myocardial contusion 355 528-30 532
 myocardial foreign body 5 5-6
 myocardial infarction 532
 pericardial foreign body 355 525-6
 pericarditis 354-5
 ruptured aorta 211 355 528
 ruptured aortic cusp 530
 ruptured heart 528-9
 ruptured mitral valve 530
 spontaneous lesions 521-5
 stab wound 354 527
 sudden death 528
 Tremor 485
 Trendelenberg operation 455
 Triangle
 Fimthoven's 68 81
 supra aortic 7
 Triangular pad of fat 35
 Triaxial reference system 81
 Tricuspid atresia 250
 Tricuspid diastolic murmur 306
 Tricuspid incompetence 303-4
 catheter studies 304
 clinical features 304
 diagnosis 303
 functional 303
 hepatic pulsation 304
 in pulmonary heart disease 462
 rheumatic 303
 venous pulse 10 304
 X ray 59 304
 Tricuspid stenosis 306-7
 cirrhosis of the liver in 306
 clinical features 306
 giant a wave 170 306
 physiology 306
 venous pulse 170 306
 X ray 59 307
 Tricuspid valvulotomy 308
 Trinitrin 381
 Triple rhythm 163-6
 Tromexan 455
 Trueta's theory 421
 Truncus arteriosus
 development of 206
 pericardial stent 250
 Trypanosomiasis South American 315
 Tsutsugamushi fever 315
 Tuberculosis (see *Pulmonary tuberculosis*)
 Tuberculous pericarditis 348 353
 Tumours
 myxomatous 193 321
 pericardial 354
 rhabdomyoma 31
 sarcomatous 120 321
 Twins 103
 Twitching
 alkalotic 201
 nocturnal 193
 in syncope 123 193
 Typhoid fever 316
 Typhus scrub 35
 Urticaria
 in angina pectoris 102
 in digitalis therapy 80
 in hypertension 80
 inversion of 80 102 126
 in left ventricular hypertrophy 80
 normal 79
 in pericardial ductus 26
 in right ventricular hypertrophy 80

- U waves (contd)
 in ventricular septal defect 222
- Ulcerative endocarditis 330
- Unfolding of the aorta 39 429
- Unilateral exophthalmus 481
- Unipolar leads 68 74
- Uræmia
 in bacterial endocarditis 334 336
 pericarditis in 355
 from sodium depletion 185
 T waves 105 326
 and vaso dilatation 321
- Urea
 blood 22
 as a diuretic 185
- Urinary creatine 488
- Urine
 in bacterial endocarditis 334
 in embolic nephritis 334
 in essential hypertension 433
 examination of 21
 in heart failure 174
 in malignant hypertension 433
 in nephritic hypertension 433
 in renal infarction 334
- Urticaria
 from penicillin 338
 in rheumatic fever 63
 in serum sickness 311
 and syncope 195
- V lead 68 74
- V wave 19 170
 and mitral diastolic murmur 286
 normal 19
 and opening snap 286
 and third heart sound 165
 in tricuspid incompetence 170 304
- Vagal influence
 on auricular fibrillation 531
 on auricular flutter 142
 in cardiac standstill 115 193
 in depressor reflex 113
 on focal myocardial necrosis 322
 on heart block 118 123
 in nodal rhythm 117
 on pace maker 109
 in pulmonary embolism 449
 in sino auricular block 115
 in sinus arrhythmia 109
 in sinus bradycardia 113
 in sudden death 528
 in syncope 194-6 197
- Vagal paralysis 110
- Valsalva's experiment 224
- Valve amplifying oscillograph 67
- Valves (see Aortic Mitral etc)
 auscultatory sites of 9
 in bacterial endocarditis 330
 calcified 61
- Valves (contd)
 congenital lesions of 21- 229-44 250
 in dissecting aneurysm 512
 position of 63 242
 in rheumatic carditis 257-8
 rheumatic lesions of 280-307
 rupture of 530
 syphilitic 364
 vascularity of 257
- Valvulitis (rheumatic) 257-8
- Van Slyke's apparatus 16
- Varicose veins and pulmonary embolism 443
- Vascular murmurs (see Murmurs) 208 487
- Vaso constriction
 in acute nephritis 417
 in congestive failure 178
 in essential hypertension 420
 in hæmorrhage 195
 in hyperventilation 61
 in left ventricular failure 160
 in massive pulmonary embolism 447
 in mitral stenosis 285
 in pericardial effusion 348
 in Pick's disease 349
 pulmonary 462
 in renal hypertension 418
- Vaso dilatation
 in anaemia 503
 in anoxic cor pulmonale 465
 in aortic incompetence 295 297
 in arteriovenous aneurysm 514
 in beri beri 516
 cholinergic 200 516
 in hepatic failure 517
 histamine 200
 in hyperkinetic circulatory states 50
 menopausal 200
 in syncope 194-6
 thyrotoxic 487
- Vaso motor collapse
 in cor pulmonale 465
 in diphtheria 311
 in syncope 195
 toxic 314
- Vaso vagal syncope 194-200
- Vector cardiac 80-2
- Vectorcardiogram 83
- Vegetative endocarditis 330
- Veins (see Venous)
- Vena cava (see I I C or S I C)
- Venesection 179
 bloodless 195
 in congestive failure 179
 in hypertension 437
 in left ventricular failure 190
 in pulmonary heart disease 474
 and syncope 195
- Veno arterial shunt (in)
 Fallot's tetralogy 236

- Barium enema in tumors of small intestine 265
 in ulcerative colitis 355
 in volvulus of small intestine 262
 meal study 61 65
 findings at various time intervals 62 64
 of duodenum 62
 of esophagus 61
 of stomach 61
 primary use 65
 Bed rest in peptic ulcer 140
 Beef tapeworm 556 560
 Belching diagnostic significance of 24
 Belladonna in peptic ulcer 176
 Benign tumors of stomach 27
 ulcer of lesser curvature roentgen views 203
 Beta alpha phenyl propionic acid in cholecystography 66
 Biliary tract duodenal drainage 54
 Bile ducts cancer 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 diseases of 485 505
 new growth differentiated from cancer of pancreas 538
 stone differentiated from cancer of pancreas 538
 stricture 494 499
 etiology 497
 pathology 498
 symptoms 498
 treatment 499
 Bile pigment enterohepatic circulation of 408
 metabolism abnormalities in hemolytic jaundice 409
 in obstructive jaundice 407
 Bilharziasis *See* Schistosomiasis
 Biliary cirrhosis hypertrophic 473
 xanthomatous skin signs 48
 colic 490
 obstruction intrahepatic 473
 stasis in infectious hepatitis 433
 tract duodenal drainage 52
 in cholelithiasis 493
 Billousness 318
 Bilirubin intensity of jaundice studied by 406
 metabolism in liver 406
 Bilirubinemia increase in obstructive jaundice 417
 Biopsy of skin in diagnosis of hemochromatosis 471
 Bland diet standard 149
 Blastogenic cysts of pancreas 516
 Bleeding in peptic ulcer 155
 Blood calcium level in acute pancreatitis 512
 chemical changes in pyloric obstruction from peptic ulcer 206
 cholesterol in obstructive jaundice 418
 diseases digestive aspects of 599
 donors excluded in infectious hepatitis 442
 findings in cancer of pancreas 531
 in infectious hepatitis 439
 flukes *See* Schistosomes
 transfusion in hemorrhage from peptic ulcer 195
 volume in hemorrhage from peptic ulcer 192
 Bodily systems relative involvement of various 26
 Body habitus 72 74
 relation to clinical conditions table 73
 types illustrated 72
 Brain lesions in peptic ulcer 147
 tissue changes in tumors of islands of Langerhans 524
 Breath odor diagnostic significance 49
 Bromsulfalein test 411
 Bulimia 128
- C**
- Calcium deficiency in steatorrhea effects 495
 gluconate in intestinal tuberculosis 273
 Calculi salivary 91
 Cancer of appendix 391
 of bile ducts 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 of body of pancreas signs and symptoms table 399
 of bowel advances in therapy 81
 of cecum roentgen view and sketch 344
 spread 372
 of colon 369 381
 biopsy for diagnosis 345
 classification 340

- Cancer of colon—Cont'd
 colloid pathology 371
 diagnosis 373 378
 distribution sketch 370
 etiology 369
 incidence 369
 pathology 370
 polypoid 370
 prognosis 379
 roentgen study 375 378
 scirrhus pathology 370
 spread 371
 symptoms 373
 treatment 378 81
- of duodenum 246
- of esophagus 107 111
 roentgen view 110
 hiatus hernia differentiated from 121
- of gallbladder 503
- of gastrointestinal tract ana-
 tomic distribution table
 71
- of head of pancreas signs and
 symptoms table 527
- of liver 477 481
 diagnosis 481
 etiology 477
 geographic distribution 478
 table 479
 laboratory findings 480
 pathology 477
 prognosis 481
 symptoms 479
 treatment 481
- of pancreas 575 5 9
 ascites 531
 blood signs 531
 creatorrhea as symptom 5 4
 diagnosis 5 6-5 8
 digestive symptoms 530
 duration 31
 fecal urobilinogen 534
 feces characteristics 5 1
 gallbladder distention 530
 glucose tolerance test 531
 head and body signs and
 symptoms compared 5 8
 incidence 575
 jaundice as symptom 5 9
 diagram of mechanism 53
 laboratory studies 504
 laboratory findings 531
 liver enlargement 5 0
 metastasis routes 526
 pain 5 8
 pancreatic ferments study of
 534
 pathology 5 6-5 7
- Cancer of pancreas—Cont'd
 radical operations for sketch
 537
 roentgen examination 534
 serum lipase in 5 4
 sites of metastasis and invasion
 table 5 5
 symptoms 577 579
 treatment 578
 weight loss 5 7
- of papilla of Vater differentiated
 from cancer of bile ducts
 500
- of rectum 369 381
 classification 370
 colloid pathology 370
 diagnosis 373 378
 distribution sketch 370
 etiology 369
 incidence 369
 obstructing view facing 60
 pathology 370
 polypoid pathology 370
 primary adenocarcinoma of
 liver from 372
 primary cancer of cervix uteri
 380
 prognosis 379
 scirrhus pathology 370
 spread 372
 symptoms 377
 treatment 378 81
- of stomach 417 240
 achlorhydria associated with
 182
 atrophic gastritis and 217
 diagnosis 2 7
 differentiated from peptic
 ulcer 167
 table 200
 dyspepsia as symptom 221
 fungating 18
 gastric analysis in 224
 incidence 217
 table 218
 operative mortality 229
 organs involved in spreading
 7 0
 pathogenesis 217
 pathology 218 220
 polypoid roentgen views 223
 prognosis 227
 roentgenologic diagnosis 224
 views 226
 specimen 2 8
 spreading 219
 symptoms 0
 treatment 279
 types schematic drawings
 ulcerated 19

- Barium enema in tumors of small intestine 265
 in ulcerative colitis 355
 in volvulus of small intestine 262
 meal study 61 65
 findings at various time intervals 62 64
 of duodenum 62
 of esophagus 61
 of stomach 61
 primary use 65
 Bed rest in peptic ulcer 170
 Beef tapeworm 556 560
 Belching diagnostic significance of 24
 Belladonna in peptic ulcer 146
 Benign tumors of stomach 227
 ulcer of lesser curvature roentgen views 203
 Beta alpha phenyl propionic acid in cholecystography 66
 Biliary tract duodenal drainage 52
 Bile ducts cancer 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 diseases of 485 503
 new growth differentiated from cancer of pancreas 538
 stone differentiated from cancer of pancreas 538
 stricture 497 499
 etiology 497
 pathology 498
 symptoms 498
 treatment 499
 Bile pigment enterohepatic circulation of 403
 metabolism abnormalities in hemolytic jaundice 409
 in obstructive jaundice 407
 Bilharziasis *See* Schistosomiasis
 Biliary cirrhosis hypertrophic 473
 xanthomatous skin signs 28
 colic 490
 obstruction intrahepatic 473
 stasis in infectious hepatitis 433
 tract duodenal drainage 52
 in cholelithiasis 493
 Biliousness 318
 Bilirubin intensity of jaundice studied by 406
 metabolism in liver 406
 Bilirubinemia increase in obstructive jaundice 417
 Biopsy of skin in diagnosis of hemochromatosis 471
 Bland diet standard 129
 Blastogenic cysts of pancreas 516
 Bleeding in peptic ulcer 155
 Blood calcium level in acute pancreatitis 512
 chemical changes in pyloric obstruction from peptic ulcer 206
 cholesterol in obstructive jaundice 418
 diseases digestive aspects of 599
 donors excluded in infectious hepatitis 442
 findings in cancer of pancreas 531
 in infectious hepatitis 439
 flukes *See* Schistosomes
 transfusion in hemorrhage from peptic ulcer 195
 volume in hemorrhage from peptic ulcer 192
 Bodily systems relative involvement of various 46
 Body habitus 72 74
 relation to clinical conditions table 43
 types illustrated 72
 Brain lesions in peptic ulcer 147
 tissue changes in tumors of islands of Langerhans 522
 Breath odor diagnostic significance 29
 Bromsulfalein test 411
 Bulimia 128
- C**
- Calcium deficiency in steatorrhea effects 295
 gluconate in intestinal tuberculosis 273
 Calcium salivary 91
 Cancer of appendix 391
 of bile ducts 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 of body of pancreas signs and symptoms table 399
 of bowel advances in therapy 381
 of cecum roentgen view and sketch 374
 spread 372
 of colon 369 381
 biopsy for diagnosis 375
 classification 370

- Cancer of colon—Cont d
 colloid pathology 371
 diagnosis 373 378
 distribution sketch 3,0
 etiology 369
 incidence 369
 pathology 3,0
 polypoid 3,0
 prognosis 379
 roentgen study 3,5 378
 scirrhus pathology 3,0
 spread 371
 symptoms 373
 treatment 378 381
- of duodenum 246
- of esophagus 107 111
 roentgen view 110
 hiatus hernia differentiated from 1 1
- of gallbladder 503
- of gastrointestinal tract ana-
 tomic distribution table
 221
- of head of pancreas signs and
 symptoms table 5 7
- of liver 477 481
 diagnosis 481
 etiology 477
 geographic distribution 4,8
 table 479
 laboratory findings 480
 pathology 477
 prognosis 481
 symptoms 479
 treatment 481
- of pancreas 525-539
 ascites 531
 blood signs 531
 creatorrhea as symptom 534
 diagnosis 5 6 538
 digestive symptoms 530
 duration 531
 fecal urobilinogen 534
 feces characteristics 531
 gallbladder distention 530
 glucose tolerance test 531
 head and body signs and
 symptoms compared 5 8
 incidence 5 5
 jaundice as symptom 5 9
 diagram of mechanism 53°
 laboratory studies 534
 laboratory findings 531
 liver enlargement 5 0
 metastasis routes 526
 pain 5 8
 pancreatic ferments study of
 534
 pathology 526 5 7
- Cancer of pancreas—Cont d
 radical operations for sketch
 537
 roentgen examination 534
 serum lipase in 534
 sites of metastasis and invasion
 table 5°5
 symptoms 527 5,3
 treatment 538
 weight loss 527
- of papilla of Vater differentiated
 from cancer of bile ducts
 500
- of rectum 369 381
 classification 370
 colloid pathology 370
 diagnosis 373 3,8
 distribution sketch 370
 etiology 369
 incidence 369
 obstructing view facing 60
 pathology 370
 polypoid pathology 3,0
 primary adenocarcinoma of
 liver from 372
 primary cancer of cervix uteri
 380
 prognosis 379
 scirrhus pathology 370
 spread 372
 symptoms 373
 treatment 378 381
- of stomach 217 240
 achlorhydria associated with
 182
 atrophic gastritis and 21,
 diagnosis 22°
 differentiated from peptic
 ulcer 167
 table 65
 dyspepsia as symptom 21
 fungating 18
 gastric analysis in 224
 incidence 17
 table 218
 operative mortality 29
 organs involved in spreading
 2 0
 pathogenesis 217
 pathology 218 20
 polypoid roentgen views 223
 prognosis 227
 roentgenologic diagnosis 244
 views 276
 specimen 228
 spreading 219
 symptoms 2°0
 treatment 229
 types schematic drawings 2
 ulcerated 19

- Barium enema in tumors of small intestine 265
 in ulcerative colitis 355
 in volvulus of small intestine 262
 meal study 61 65
 findings at various time intervals 62 64
 of duodenum 62
 of esophagus 61
 of stomach 61
 primary use 65
 Bed rest in peptic ulcer 170
 Beef tapeworm 556 560
 Belching diagnostic significance of 24
 Belladonna in peptic ulcer 176
 Benign tumors of stomach 227
 ulcer of lesser curvature roentgen views 203
 Beta alpha phenyl propionic acid in cholecystography 66
 Biliary tract duodenal drainage 52
 Bile ducts cancer 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 diseases of 485 505
 new growth differentiated from cancer of pancreas 538
 stone differentiated from cancer of pancreas 538
 stricture 497 499
 etiology 497
 pathology 498
 symptoms 498
 treatment 499
 Bile pigment enterohepatic circulation of 408
 metabolism abnormalities in hemolytic jaundice 409
 in obstructive jaundice 407
 Bilharziasis *See* Schistosomiasis
 Biliary cirrhosis hypertrophic 473
 xanthomatous skin signs 28
 colic 490
 obstruction intrahepatic 473
 stasis in infectious hepatitis 433
 tract duodenal drainage 52
 in cholelithiasis 493
 Billousness 318
 Bilirubin intensity of jaundice studied by 406
 metabolism in liver 406
 Bilirubinemia increase in obstructive jaundice 417
 Biopsy of skin in diagnosis of hemochromatosis 471
 Bland diet standard 129
 Blastogenic cysts of pancreas 516
 Bleeding in peptic ulcer 155
 Blood calcium level in acute pancreatitis 512
 chemical changes in pyloric obstruction from peptic ulcer 206
 cholesterol in obstructive jaundice 418
 diseases digestive aspects of 599
 donors excluded in infectious hepatitis 442
 findings in cancer of pancreas 531
 in infectious hepatitis 439
 flukes *See* Schistosomes
 transfusion in hemorrhage from peptic ulcer 195
 volume in hemorrhage from peptic ulcer 192
 Bodily systems relative involvement of various 76
 Body habitus 72 74
 relation to clinical conditions table 73
 types illustrated 72
 Brain lesions in peptic ulcer 147
 tissue changes in tumors of islands of Langerhans 522
 Breath odor diagnostic significance 29
 Bromsulfalein test 411
 Bulimia 128
- C**
- Calcium deficiency in steatorrhea effects 795
 gluconate in intestinal tuberculosis 273
 Calculi salivary 91
 Cancer of appendix 391
 of bile ducts 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 of body of pancreas signs and symptoms table 5 9
 of bowel advances in therapy 381
 of cecum roentgen view and sketch 314
 spread 372
 of colon 369 81
 biopsy for diagnosis 375
 classification 310

Cancer of colon—Cont'd

- colloid pathology 371
- diagnosis 3,3 3,8
- distribution sketch 3,0
- etiology 69
- incidence 369
- pathology 370
- polypoid 3,0
- prognosis 379
- roentgen study 375 18
- scirrhous pathology 3,0
- spread 371
- symptoms 373
- treatment 3,8 81
- of duodenum 46
- of esophagus 107 111
 - roentgen view 110
 - hiatus hernia differentiated from 121
- of gallbladder 303
- of gastrointestinal tract and
 - tonic distribution table 371
- of head of pancreas signs and
 - symptoms table 37
- of liver 47 481
 - diagnosis 481
 - etiology 4,7
 - geographic distribution 4,8
 - table 479
 - laboratory findings 480
 - pathology 4,1
 - prognosis 481
 - symptoms 4,9
 - treatment 481
- of pancreas 525 9
 - ascites 531
 - blood sign 531
 - creatorrhea as symptom 534
 - diagnosis 6538
 - digestive symptoms 530
 - duration 531
 - fecal urobilinogen 534
 - feces characteristics 51
 - gallbladder distention 0
 - glucose tolerance test 51
 - head and body signs and
 - symptoms compared 528
 - incidence 5
 - jaundice as symptom 59
 - diagram of mechanism 59
 - laboratory studies 34
 - laboratory findings 531
 - liver enlargement 50
 - metastasis routes 6
 - pain 8
 - pancreatic ferments study of 54
 - pathology 5 6-5,7

Cancer of pancreas—Cont'd

- radical operations for sketch 337
- roentgen examination 534
- serum lipase in 534
- sites of metastasis and invasion,
 - table 525
- symptoms 527 539
- treatment 538
- weight loss 5,7
- of papilla of Vater differentiated from cancer of bile ducts 300
- of rectum 369 381
 - classification 3,0
 - colloid pathology 370
 - diagnosis 373 3,8
 - distribution sketch 370
 - etiology 369
 - incidence 369
 - obstructing view facing 60
 - pathology 3,0
 - polypoid pathology 3,0
 - primary adenocarcinoma of liver from 3,2
 - primary cancer of cervix uteri 30
 - prognosis 379
 - scirrhous pathology 3,0
 - spread 3,2
 - symptoms 3,7
 - treatment, 3,8 381
- of stomach 371 40
 - achlorhydria associated with 12
 - atrophic gastritis and 217
 - diagnosis 2,2
 - differentiated from peptic ulcer 167
 - table 03
 - dyspepsia as symptom 21
 - fungating 18
 - gastric analysis in 2,4
 - incidence 17
 - table 218
 - operative mortality 229
 - organs involved in spreading 220
 - pathogenesis 217
 - pathology 218 2,0
 - polypoid roentgen views 22
 - prognosis 37
 - roentgenologic diagnosis 24
 - views 2,6
 - specimen 8
 - spreading 19
 - symptoms 220
 - treatment 9
 - types schematic drawings 3,2
 - ulcerated 219

- Barium enema in tumors of small intestine 265
 in ulcerative colitis 355
 in volvulus of small intestine 262
 meal study 61 65
 findings at various time intervals 62 64
 of duodenum 62
 of esophagus 61
 of stomach 61
 primary use 65
- Bed rest in peptic ulcer 170
- Beef tapeworm 556 560
- Belching diagnostic significance of 24
- Belladonna in peptic ulcer 176
- Benign tumors of stomach 227
 ulcer of lesser curvature roentgen views 203
- Beta alpha phenyl propionic acid in cholecystography 66
- Biliary tract duodenal drainage 52
- Bile ducts cancer 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 diseases of 485 505
 new growth differentiated from cancer of pancreas 538
 stone differentiated from cancer of pancreas 538
 stricture 497 499
 etiology 497
 pathology 498
 symptoms 498
 treatment 499
- Bile pigment enterohepatic circulation of 408
 metabolism abnormalities in hemolytic jaundice 409
 in obstructive jaundice 407
- Bilharziasis *See* Schistosomiasis
- Biliary cirrhosis hypertrophic 473
 xanthomatous skin signs 98
 colic 190
 obstruction intrahepatic 473
 stasis in infectious hepatitis 433
 tract duodenal drainage 52
 in cholelithiasis 493
- Biliousness 318
- Bilirubin intensity of jaundice studied by 406
 metabolism in liver 406
- Bilirubinemia increase in obstructive jaundice 417
- Biopsy of skin in diagnosis of hemochromatosis 411
- Bland diet standard 129
- Blastogenic cysts of pancreas 516
- Bleeding in peptic ulcer 155
- Blood calcium level in acute pancreatitis 512
 chemical changes in pyloric obstruction from peptic ulcer 206
 cholesterol in obstructive jaundice 418
 diseases digestive aspects of 599
 donors excluded in infectious hepatitis 442
 findings in cancer of pancreas 531
 in infectious hepatitis 439
 flukes *See* Schistosomes
 transfusion in hemorrhage from peptic ulcer 195
 volume in hemorrhage from peptic ulcer 192
- Bodily systems relative involvement of various 26
- Body habitus 72 74
 relation to clinical conditions table 73
 types illustrated 72
- Brain lesions in peptic ulcer 147
 tissue changes in tumors of islands of Langerhans 522
- Breath odor diagnostic significance 29
- Bromsulfalein test 411
- Bulimia 128

C

- Calcium deficiency in steatorrhea effects 295
 gluconate in intestinal tuberculosis 273
- Calculi salivary 91
- Cancer of appendix 391
 of bile ducts 499 501
 diagnosis 500
 incidence 499
 pathology 499
 symptoms 500
 treatment 501
 of body of pancreas signs and symptoms table 39
 of bowel advances in therapy 81
 of cecum roentgen view and sketch 374
 spread 372
 of colon 369 381
 biopsy for diagnosis 375
 classification 310

Cancer of colon—Cont'd

- colloid pathology 3.1
- diagnosis 3.3-3.8
- distribution sketch 3.0
- etiology 3.69
- incidence 3.69
- pathology 3.0
- polypoid 3.0
- prognosis 3.9
- roentgen study 3.5-3.8
- scirrhus pathology 3.0
- spread 3.1
- symptoms 3.3
- treatment 3.8-81
- of duodenum 246
- of esophagus 107-111
 - roentgen view 110
 - hiatus hernia differentiated from 1.1
- of gallbladder 03
- of gastrointestinal tract anatomic distribution, table 221
- of head of pancreas signs and symptoms table 5.7
- of liver 477-481
 - diagnosis 481
 - etiology 477
 - geographic distribution 4.8
 - table 479
 - laboratory findings 480
 - pathology 477
 - prognosis 481
 - symptoms 4.9
 - treatment 481
- of pancreas 5.5-9
 - ascites 3.1
 - blood signs 3.1
 - creatorrhea as symptom 3.4
 - diagnosis 536-8
 - digestive symptoms 5.0
 - duration 31
 - fecal urobilinogen 534
 - feces characteristics 531
 - gallbladder distention 0
 - glucose tolerance test 531
 - head and body signs and symptoms compared 5.8
 - incidence 5.0
 - jaundice as symptom 3.9
 - diagram of mechanism 5.3
 - laboratory studies 531
 - laboratory findings 531
 - liver enlargement 5.0
 - metastasis routes 526
 - pain 5.8
 - pancreatic ferments study of 534
 - pathology 526-7

Cancer of pancreas—Cont'd

- radical operations for sketch 53.1
- roentgen examination 534
- serum lipase in 534
- sites of metastasis and invasion
 - table 5.5
- symptoms 527-539
- treatment 5.8
- weight loss 5.7
- of papilla of Vater differentiated from cancer of bile ducts 500
- of rectum 369-381
 - classification 3.0
 - colloid pathology 3.0
 - diagnosis 3.3-3.8
 - distribution sketch 3.0
 - etiology 69
 - incidence 69
 - obstructing view facing 60
 - pathology 3.0
 - polypoid pathology 3.0
 - primary adenocarcinoma of liver from 3.1
 - primary cancer of cervix uteri 380
 - prognosis 379
 - scirrhus pathology 3.0
 - spread 372
 - symptoms 373
 - treatment, 3.8-81
- of stomach 17-40
 - achlorhydria associated with 13
 - atrophic gastritis and 21.1
 - diagnosis 22.1
 - differentiated from peptic ulcer 167
 - table 105
 - dyspepsia as symptom 11.1
 - fungating 18
 - gastric analysis in 2.4
 - incidence 217
 - table 218
 - operative mortality 2.9
 - organs involved in spreading 20
 - pathogenesis 217
 - pathology 218-20
 - polypoid roentgen views 2.3
 - prognosis 2.7
 - roentgenologic diagnosis 2.4
 - views 6
 - specimen 2.8
 - spreading 219
 - symptoms 220
 - treatment 9
 - types schematic drawings 19
 - ulcerated 19

- Canker sores allergic aspect 622
treatment 81
- Carbirsone in amebiasis 552
- Carbohydrate metabolism in liver 409
- Carcinoids of appendix 391
- Carcinoma *See* Cancer
- Cardiac cirrhosis 450 452
diagnosis 452
pathology 450
symptoms 451
sphincter achalasia in esophageal spasm 97
- Cardiospasm 95 99 126 *See also* Esophagus spasm of
functional theories of 95 96
- Cardiovascular diseases digestive disorders in 596
- Cathartics abuse of constipation caused by 323
- Cecocolon hyperfixation of 10
- Cecum cancer of roentgen view and sketch 374
spread 37
high 309
low 307
symptoms 308
treatment 308
nondescent of 308
- Celiac disease *See* Steatorrhea idiopathic
- Central nervous system manifestations of schistosomiasis japonica 516
- Cephalin cholesterol flocculation test 404
- Cerebrospinal syphilis digestive symptoms 593
- Cestodes 556 563
mode of infestation 556
- Cheilitis acute allergic aspect of 622
- Cheilosis acute from riboflavin deficiency illustration 83
- Chiari's syndrome 103 404
diagnosis 405
etiology 454
pathology 454
prognosis 455
symptoms 454
treatment 455
- Chinofon in amebiasis 352
- Chloride depletion in pyloric obstruction from peptic ulcer 306
- Cholecystectomy 496
persistence of symptoms following 437
- Cholecystitis 480 488
complications 487
diagnosis 487
differentiated from acute appendicitis 389
etiology 485
pathogenesis 485
pathology 486
symptoms 487
treatment 488
- Cholecystography 66 67 491
preparatory diet 67
technic 66
- Choledochodochorrhaphy 499
- Choledochoduodenostomy 199
- Choledocholithiasis 501 503
diagnosis 502
incidence 501
pathogenesis and pathology 501
pathology 416
prognosis 503
symptoms 503
treatment 502
- Cholelithiasis 488 497
complicating hemolytic jaundice 421
diagnosis 490
diet in 494
differential diagnosis 49
incidence 488
pathogenesis 488
pathology 489
symptoms 490
treatment 494
- Cholera sicca 215
- Cholesterol metabolism in liver 403
- Cholesterosis of gallbladder 486
- Cirrhosis atrophic 46 471 *See also* Cirrhosis portal
pathology 463
biliary hypertrophic 410
xanthomatous skin signs 98
cancer of liver and relationship 479
cardiac 450 452 *See also* Cardiac cirrhosis
due to schistosomiasis section view of liver 564
- Laennec's 462 411
portal 46 411
cancer of liver and section view 418
collateral circulation in 466
definition 462
diagnosis 469
laboratory findings 468
liver function tests 469
omentopexy for 411
section in view 463
surgical treatment 471

- Cirrhosis portal—Cont d
 symptoms 466
 treatment 469
 varicose veins at lower end of
 esophagus view 464
 xanthomatous biliary skin signs
 8
- Classification of digestive disorders
 17
- Clinical conditions relation to body
 habitus table 3
- Lila orclis sin nsis* 18
- Coated tongue diagnostic signifi-
 cance 23
- Colectomy subtotal in Hirsch
 sprungs disease 31
- Colic biliary 430
- Colicky pain diagnostic signifi-
 cance 26
- Colitis differentiation from peptic
 ulcer 168
mucous 2839
 diagnosis 39
 treatment 39
 simple See Colon unstable
 ulcerative 3426
 allergic aspect 64
 complications 24251
 incidence table 248
 definition 343
 diagnosis 3251
 diet 228
 distal ileitis and sketch 34
 emotional factors 246
 extreme emaciation and illus-
 trated 349
 five years duration roentgen
 views 254
 ileostomy in 261
 ileum involved illustrated 45
 improvement affecting leg
 ulcers 21
 mild treatment 227
 pathologic physiology 247
 pathology 346
 penicillin in 360
 perforation and illustrated 344
 postoperative treatment 262
 psychogenesis 613
 psychotherapy 329 614
 roentgen diagnosis 2
 stricture and roentgen view
 20
 sulfonamides in 60
 symptoms 221322
 treatment 57362
 psychotherapy in 359 614
 supportive 37
 surgical 61
- Collateral circulation in portal
 obstruction 466
 diagram 465
- Colloid carcinoma of colon and
 rectum pathology 340
- Colloidal gold reaction in liver
 function 413
- Colon adenomas 67
 adenomatosis hereditary 268
 anomalies of 24312
 constipation due to 313
 cancer of 69281 See also under
 Cancer
 consciousness 323
 diseases of 204236
 disorders of pain in 20
 diverticulitis and diverticulosis
 6367
 complications 22
 diagnosis 64
 etiology 362
 pathology 262
 prognosis 64
 roentgen views 63366
 specimen 362
 symptoms 264
 treatment 367
 granuloma nonspecific 342
 hereditary polyposis 368
 irritable See Colon unstable
 napkin ring carcinoma roentgen
 view 376
 nonrotation of 209
 schematic drawing 310
 polyps 368
 proximal fixation of 202
 redundant 24307
 constipation due to 317
 diagnosis 306
 dyschezia in 306
 schematic drawings 302 306
 symptoms 204
 treatment 206
 spastic See Colon unstable
 tuberculosis of illustrated 69
 tumors 36, 381
 benign 67268
 unstable 2328
 definition 322
 diagnosis 224
 differential 322
 differentiation from peptic
 ulcer 168
 etiology 32
 findings with opaque enema 322
 incidence 3
 psychic factors in 323
 psychotherapy for 323
 roentgen diagnosis 325

- Canker sores allergic aspect 62^o
treatment 81
- Carbarsone in amebiasis 552
- Carbohydrate metabolism in liver 402
- Carcinoids of appendix 391
- Carcinoma *See* Cancer
- Cardiac cirrhosis 450 452
diagnosis 452
pathology 450
symptoms 451
sphincter achalasia in esophageal spasm 97
- Cardiospasm 95-99 126 *See also* Esophagus spasm of
functional theories of 95 96
- Cardiovascular diseases digestive disorders in 596
- Cathartics abuse of constipation caused by 32_o
- Cecocolon hyperfixation of 310
- Cecum cancer of roentgen view and sketch 314
spread 372
high 309
low 307
symptoms 308
treatment 308
nondescent of 308
- Celiac disease *See* Steatorrhea idiopathic
- Central nervous system manifestations of schistosomiasis japonica 576
- Cephalin cholesterol flocculation test 404
- Cerebrospinal syphilis digestive symptoms 595
- Cestodes 5 6 563
mode of infestation 556
- Cheilitis acute allergic aspect of 622
- Cheilosis acute from riboflavin deficiency illustration 8_o
- Chfari's syndrome 10_o 44
diagnosis 45_o
etiology 454
pathology 454
prognosis 455
symptoms 454
treatment 455
- Chiniofon in amebiasis 5_o^o
- Chloride depletion in pyloric obstruction from peptic ulcer 206
- Cholecystectomy 496
persistence of symptoms following 497
- Cholecystitis 48_o 488
complications 487
diagnosis 487
differentiated from acute appendicitis 89
etiology 485
pathogenesis 485
pathology 486
symptoms 487
treatment 488
- Cholecystography 66 67 491
preparatory diet 67
technic 66
- Choledochodochorrhaphy 499
- Choledochoduodenostomy 499
- Choledocholithiasis 501 503
diagnosis 50^o
incidence 501
pathogenesis and pathology 501
pathology 416
prognosis 503
symptoms 503
treatment 502
- Cholelithiasis 488 497
complicating hemolytic jaundice 41
diagnosis 490
diet in 494
differential diagnosis 493
incidence 488
pathogenesis 488
pathology 489
symptoms 490
treatment 494
- Cholera sicca 75
- Cholesterol metabolism in liver 40_o
- Cholesterosis of gallbladder 486
- Cirrhosis atrophic 462 471 *See also* Cirrhosis portal
pathology 463
biliary hypertrophic 473
xanthomatous skin signs 28
cancer of liver and relationship 479
cardiac 450 45^o *See also* Cardiac cirrhosis
due to schistosomiasis section view of liver 567
- Laennec's 46^o 471
portal 462 471
cancer of liver and section view 478
collateral circulation in 466
definition 462
diagnosis 469
laboratory findings 468
liver function tests 469
omentopexy for 471
section in view 463
surgical treatment 471

- Cirrhosis portal--Cont'd
 symptoms 466
 treatment 469
 varicose veins at lower end of
 esophagus view 464
 xanthomatous biliary skin signs
 28
- Classification of digestive disorders
 17
- Clinical conditions relation to body
 habitus table 3
- Clonorchis sinensis* 578
- Coated tongue diagnostic signifi-
 cance 25
- Colectomy subtotal in Hirsch-
 sprungs disease 319
- Colic biliary 490
- Colicky pain diagnostic signifi-
 cance 26
- Colitis differentiation from peptic
 ulcer 168
- mucous 383 9
 diagnosis 39
 treatment 39
- simple See Colon unstable
- ulcerative 436
 allergic aspect 64
 complications 473 1
 incidence table 348
 definition 343
 diagnosis 355-57
 diet 58
 distal ileitis and sketch 342
 emotional factors 46
 extreme emaciation and illus-
 trated 349
 five years duration roentgen
 views 54
 ileostomy in 61
 ileum involved illustrated 45
 improvement affecting leg
 ulcers 51
 mild treatment 57
 pathologic physiology 41
 pathology 346
 penicillin in 60
 perforation and illustrated 344
 postoperative treatment 362
 psychogenesis 613
 psychotherapy 9 614
 roentgen diagnosis 35
 stricture and roentgen view
 30
 sulfonamides in 60
 symptoms 1 55
 treatment 35, 52
 psychotherapy in 359 614
 supportive 57
 surgical 61
- Collateral circulation in portal
 obstruction 466
 diagram 465
- Colloid carcinoma of colon and
 rectum pathology 30
- Colloidal gold reaction in liver
 function 413
- Colon adenomas 67
 adenomatosis hereditary 568
 anomalies of 504 312
 constipation due to 315
 cancer of 369 381 See also under
 Cancer
- consciousness 373
 diseases of 504 386
 disorders of pain in 50
 diverticulitis and diverticulosis
 562 367
 complications 63
 diagnosis 64
 etiology 365
 pathology 362
 prognosis 64
 roentgen views 365-366
 specimen 365
 symptoms 364
 treatment 367
- granuloma nonspecific 34
 hereditary polyposis 368
 irritable See Colon unstable
- napkin ring carcinoma roentgen
 view 376
- nonrotation of 59
 schematic drawing 310
- polyps 368
- proximal fixation of 503
- redundant 504 504
 constipation due to 317
 diagnosis 506
 dyschezia in 306
 schematic drawings 505 306
 symptoms 504
 treatment 506
- spastic See Colon unstable
- tuberculosis of illustrated 69
- tumors 369 381
 benign 367 68
 unstable 22 378
 definition 322
 diagnosis 324
 differential 325
 differentiation from peptic
 ulcer 168
 etiology 325
 findings with opaque enema 325
 incidence 32
 psychic factors in 33
 psychotherapy for 328
 roentgen diagnosis 35

- Colon unstable—Cont'd
 symptoms 324
 treatment 325-328
 Colonic irrigations 280
 Common channel theory of pancreatitis 599
 Compensatory mechanism effects of variations 71
 Complaint chief value in diagnosis 19
 Congenital hemolytic jaundice 1-0 42"
 megacolon 311
 Congestive heart failure digestive disorders in 596
 Constipation 313-322
 abuse of cathartics as cause of 3-3
 anomalies of colon as cause 313
 atonic 315
 treatment 321
 definition 313
 diagnosis 319
 diet as a cause of 314
 dyskinetic 316
 endocrine factors in 315
 etiology 313
 habit formation as cause of 315
 incidence 313
 mental hygiene in 321
 psychic factors in 314
 rectal 316
 varieties 317
 redundant colon and 317
 roentgen examination in 319
 spastic 316
 symptoms 318
 treatment 321
 varieties 315-318
 Constitutional inferiority 71-8
 types 17
 Contrast enema 66
 Convalescent ulcer diet 174
 Corneal ring in hepatolenticular degeneration 474
 Coronary artery disease differentiation from gallstones 493
 digestive disorders in 597
 Corrosive gastritis acute 135
 Corset abdominal for visceroptosis 76
 Crater ulcer 160
 illustrated 158 159 161
 Creatorrhea in cancer of pancreas 534
 Cryptitis anal with papillitis
 anoscopic view facing 60
 Cystography beta alpha phenyl propionic acid in 66
 Cysts benign of pancreas 516-519
See also under Pancreas
- D**
- Deficiency disorders oral manifestations of 8° 90
 Degeneration fatty of liver *See* Fatty metamorphosis of liver
 De leading in plumbism 602
 Destruction of intake factors increasing 86
 Detoxifying function of liver 41°
 Dew poison *See* Uncinaria
 Diabetes in cancer of pancreas 5-0
 Diabetic ketosis digestive symptoms 599
 Diagnostic methods history 18-27
 physical examination 27-39
 special tests 40-70
 Diagnostic significance of abdominal tenderness and pain 33-47
 of Argyll Robertson pupil 29
 of belching 24
 of coated tongue 2-3
 of feeling of fullness after eating 45
 of globus hystericus 25
 of halitosis 26
 of Horner's syndrome 29
 of inguinal metastases 3
 of lymph nodes in neck 0
 of nausea 22
 of pain 20
 of Virchow's node 31
 of vomiting 23
 Diaphragm eversion of 11-
 Diaphragmatic hernia 115-151
 types 116
 Diarrhea 75-80
 clinical classification 116
 table 277
 definition 275
 diagnosis 278
 drugs in treatment of 2-0
 etiology 216
 examination of stool 218
 general considerations 115
 in idiopathic steatorrhea 94
 incidence 276
 parts of intestine involved 216
 pathology 216
 proctoscopy in 213
 stool in 118
 symptoms 218
 treatment 219-280
Dibothriocephalus latus 560

- Diet bland 129
 constipation due to faulty 314
 in cholelithiasis 434
 in diagnosis of allergies 678
 in enteritis 89
 in hemorrhage from peptic ulcer 194
 in hepatitis 443
 in hyperinsulinism 508
 in intestinal tuberculosis 772
 in obstructive jaundice 419
 in peptic ulcer 170 179
 in plumbism 609
 in regional ileitis 292
 in steatorrhea 299
 in ulcerative colitis 358
 intestinal protein 358
 low fat low cholesterol 490
- Digestive complaints incidence
 table 6
 history chronological 6
 tract normal roentgen findings 63
- Dilation of lower end of esophagus 119
 of stomach acute 141
- Diodoquin in amebiasis 551
- Distomum crassum* 580
- Diverticulitis and diverticulosis of colon 36 367 *See also* under Colon
- Diverticulosis 77 78
 diagrammatic scheme 7
 Diverticulum Meckels 249 251
 of duodenum 244
 of esophagus 93 95
 of jejunum and ileum 248
 of stomach 114
- Dry mouth 90
- Duct of Wirsung atypical 06
- Duodenal bulb markedly deformed
 illustrated 163
 normal cycle of emptying 153
 serialgram 151
- Duodenal drainage 52 55
 diagnostic uses 52
 evaluation of findings 54
 materials and technic 50 54
 illustrated 3
 of biliary tract 59 493
 in cholelithiasis 493
- Duodenal ulcer anxiety and 619
 crater formation and serial gram 156
 pathologic features 148
 perforated differentiated from acute appendicitis 389
 roentgen views 09
- Duodenitis 246
- Duodenum anomalies of 241 44
 diagnosis 243
 incidence 241
 symptoms 243
 synoptic chart 242
 treatment 243
 varieties 41
 barium meal study of 69
 cancer of 716
 diverticulum of 244
 fistula of external 200
 second portion ulcer of 710
- Dyschezia 316
 in redundant colon 306
 varieties 317
- Dysenteries diagnosis of 249
- Dysentery amebic symptoms 547
 bacillary 33 343
 bacteriology 3 2
 definition 332
 diagnosis 338
 earliest lesion 333
 etiology 2
 incidence 339
 pathogenesis and pathology 339
 prognosis 343
 specimen views 334 3 5
 stages illustrated 337
 symptoms 336
 transition from acute to chronic form 333
 sketches 338 339 340
 treatment 340-343
- Dyskinetic constipation 316
- Dyspepsia differentiated from
 peptic ulcer 16,
 functional 125 198
 of phthisis 598
 symptom in cancer of stomach 21
- Dysphagia functional 10
 hysterical 101
 in carcinoma of esophagus 108
- E
- Ears examination of 9
- Eczema of lips allergic aspect of 670
- Edema angioneurotic 6 3
- Elimination diet for diagnosis of allergies 6 8
- Embolism digestive disorders associated with 597
 mesenteric 205
- Emetine in amebiasis 5 0
 toxic manifestations 551
- Emotional conflict revival of 605
 factors in ulcerative colitis 346
 upsets leading to gastric lesions 611

- Endamoeba histolytica* 541
 Endocrine factors in constipation 315
 Endometriosis of intestine 381
 Enema contrast 66
 opaque 65 66
 in unstable colon findings 375
 Enteritis 280 283
 diagnosis 281
 diet in 282
 drugs in 82
 etiology 280
 hyperplastic stenosing 280
 pathology 281
 symptoms 281
 treatment 282
Enterobius vermicularis 590
 Enterogastrone therapy in peptic ulcer 184
 Enterohepatic circulation of bile pigments 408
 Epigastric hernia 251
 discovery by palpation 37
 Epilepsy simulating hypoglycemia 524
 Epinephrine in allergy 630
 Esophageal hiatus hernia 116-114
 diagnosis 110
 differential diagnosis 121
 pathogenesis 120
 roentgenographic views 120
 122 123
 symptoms 120
 treatment 121 124
 medical 121
 surgical 114
 types 117
 illustrated 118
 lesions pain in 20
 varices 107 104
 diagnosis 104
 pathologic process 102 103
 treatment 104
 Esophagitis 100
 Esophagoscopy in sarcoma of stomach 271
 Esophagus 9, 112
 anomalies 9,
 barium meal study of 61
 cancer of 107 111
 diagnosis 109
 differentiation 109
 etiology and incidence 10,
 pathology 108
 prognosis 111
 roentgen view 111
 symptoms 108
 treatment 111
 congenitally short 119
 dilatation of lower end of 119
 Esophagus—Cont'd
 paralysis of 111
 pulsion diverticula 93
 diagnosis 94
 pathology 93
 roentgenographic views 94
 symptoms 93
 treatment 94 95
 spasm of 9, 99
 diagnosis 98
 etiology 95 96
 incidence 95
 pathogenesis and pathology 96 98
 roentgenographic view 97
 symptoms 98
 treatment 98 99
 stricture of 101
 traction diverticula of 95
 uncommon diseases of 111 11
 Esterification of cholesterol 406
 Evagination of diaphragm 115
 Ewald meal in test for gastric acidity 43
 relation of achlorhydria to age shown by 131
 Examination of ears 29
 of eyes 29
 of heart 32
 of lungs 32
 of mouth 21
 of neck 30
 of nose 29
 of pelvis 38
 of rectum 37
 of skin 27 9
 of tongue 20
 Excretion factors increasing 80
 Excretions oral disorders of 80 81
 Excretory function of liver 411
 Extradigestive diseases digestive symptoms in 595 607
 Eyes examination of 29
- F
- Fasciola hepatica* 579
Fasciolopsis busqi 580
 Fistling stomach aspiration of 47
 Fat absorption in obstructive jaundice 417
 accumulation prevention by diet 405
 metabolism in liver 405
 Fattening of asthenic invalid 75
 Fatty metamorphosis of liver 460 462
 hormones in 462
 Fear disguise of by socially acceptable complaints 611

- Fecal nitrogen increase in cancer of pancreas 534
- Feces *See also* Stool
bulk increase in pancreatic disorders 507
examination of 56
in amebiasis 548
in cancer of pancreas 51
in diarrhea 78
tests of 56
chemical 56
microscopic 57
- Femoral hernia 751
- Fibrosarcoma of colon and rectum pathology 310
- Fissure internal exposing sphincter fibers anoscopic view *See* 9 60
of rectum 94
- Fistula formation in regional ileitis 89
of duodenum external 70
of ileum 753
of jejunum 753
of rectum 734
of small intestine 257 206
- Fixation of proximal colon 89
- Flagellates 55 554
- Flatworms (cestodes) 5 5063
- Flatworms (trematodes) 564 531
- Fluid accumulation of in dilatation of stomach 141
- Flukes blood 564 578 *See* *if* Schistosomiasis
intestinal 780
liver 578 780
- Fluoroscopy preliminary 60
- Folic acid in sprue syndrome 300
- Food diaries in diagnosis of allergies 68
mother love and 606
pain in peptic ulcer relieved by 151
- Foreign bodies in stomach 2 6
- Formol gel test 412
- Fractional method in gastric analysis 4
- Fret test for lymphopathia venereum of intestine 274
- Fuadin in schistosomiasis japonica 517
- Fullness feeling of diagnostic significance 25
- Fundus carcinoma of roentgen views 225
- Fungating carcinoma of stomach 218
- G
Galactose tolerance test 40
- Gallbladder cancer of 503
cholesterols of 486
differentiated from peptic ulcer 167
diseases of 485-500
distended in cancer of pancreas 530
normal roentgen view 67
visualization in cholelithiasis 491
- Gallstones 488 497 *See also* Cholelithiasis
roentgen views 492
- Gas distress in constipation due to redundant colon 318
- Gastrectomy subtotal in peptic ulcer 180
serialgram 12
- Gastric *See also* Stomach
acidity Ewald test meal in 43
need for two determinations 44
pain produced by 152
significance of 44
tests for 43
analysis 40-50
conditions in which useful 40
in cancer of stomach 274
limitations 40
technic 41
cancer *See also* Cancer of stomach
achlorhydria associated with 122
contents in peptic ulcer 145
emptying delayed in peptic ulcer 149
time of some basic foods 49
function emotions in 169
histamine test for 45
Hollander's test of 46
hydrochloric acid in test of 48
insulin test for 45
motor test meal 49
neutral red in test of 41
Palmer's test 48
hyperacidity 126
hyperalgesia 17
hyperesthesia 127
irritation dietary treatment 179
medical treatment 10
syndrome of 18
lesions emotional upsets leading to 611
neutral red for check on progress 48
motility increased and diminished 126
motor function disorders of 16

- Gastric—Cont d
 pneumatosis 24
 polyps 227
 re-ection achlorhydria associated with 132
 for cancer of stomach 229
 secretions disorders of 126
 sensation disorders of 127
 subacidity 124
 test meals secretory 42
 ulcer *See also* Peptic ulcer
 gastroscopic picture character-
 istic 166
 malignant degeneration in
 202-203
 Gastritis 134-141
 acute corrosive 15
 hematogenous 135
 phlegmonous 175
 simple 134
 antral 140
 in peptic ulcer 146
 atrophic gastric carcinoma and
 140
 pathology 137
 relation to gastric cancer 217
 symptoms 139
 chronic 136-141
 achlorhydria associated with
 132
 course 140
 diagnosis 139
 etiology 136
 gastroscopy in 139
 incidence 136
 pathology 136
 symptoms 137
 treatment 140
 classification 134
 hematogenous acute 135
 hypertrophic pathology 137
 symptoms 139
 peptic ulcer and 140
 postoperative 140
 superficial pathology 136
 symptoms 137
 Gastroduodenal *See* Peptic
 Gastroenterostomy in peptic ulcer
 sequelae of 180
 Gastrointestinal allergy 618-633
 See also under Allergy
 hemorrhage *See under*
 Hemorrhage
 symptoms intensification by
 threat to security 610
 system emphasis on in child
 training 607
 tract cancer of anatomic distri-
 bution 221
 susceptibility to acid injury 146
- Gastrojejunal ulcer 211
 Gastropptosis 114
 Gastroscopic findings in
 achlorhydria 131
 in hemorrhage from peptic
 ulcer 193
 Gastroscopy in chronic gastritis 130
 Gee Herter's disease 293
 Genitourinary diseases digestive
 symptoms 600
 Gentian violet in strongyloidiasis
 392
 Giardia infection 552
 atabrine in 553
 Gingivitis pathologic process 8
 treatment 8
 Globus hystericus diagnostic
 significance 75
 Glossitis chronic 86-90
 areas of tongue affected 87
 illustrations of stages and
 types 88-89
 pathogenesis 87
 treatment 87
 diagnostic significance 30
 Glossodynia 90
 Gluconeogenesis and glycogenolysis
 in liver 402
 Glucose tolerance test in cancer of
 pancreas 531
 Glycosuria in cancer of pancreas
 530
 Granuloma nonspecific of colon
 sketch 344
 Gray Turner sign in acute
 pancreatitis 510
 Ground itch *See* Uncinaria
- H
- Habit formation in constipation 315
 Habitus 74
 Hair balls in stomach 236
 Halitosis diagnostic significance 26
 treatment 81
 Hanger test 404
 Healing in tuberculosis of intestine
 69
 Heart disorders diagnostic
 significance
 examination of 32
 Heartburn 24
 Hematemesis in cancer of pancreas
 530
 in peptic ulcer 155
 Hematogenous gastritis acute 135
 Hemochromatosis 411
 in pancreatitis 514
 skin signs 28

- Hemolytic jaundice 415 420 42
 bile pigment metabolism in 407
 abnormalities 409
 definition 420
 diagnosis 422
 etiology 420
 laboratory findings 421
 pathology 40
 symptoms 40
 treatment 422
- Hemorrhage from peptic ulcer
 191 199
 anatomic lesion how
 demonstrated 193
 blood transfusion 195
 blood volume determination
 192
 diagnosis 192
 diet 194
 drugs in treatment 196
 gastroscopy 193
 mortality statistics 199
 prognosis 197
 recovery criteria 198
 surgical treatment 197
 treatment 194 199
- Hemorrhagic erosions gastric 147
- Hemorrhoids 399
 acute thrombotic internal procto-
 scopic view *facing* 60
 chronic internal proctoscopic
 view *facing* 60
 treatment 400
- Hepatic cell dysfunction in
 obstructive jaundice 417
 dysfunction constitutional 42
 stage of infectious jaundice 446
- Hepatitis acute differentiation
 from gallstones 493
 arsenical 448
 cholangiolitic 442
 chronic course 441
 diagnosis 441
 laboratory findings 441
 menopausal 443
 symptoms 440
 fatal 433
 infectious 464 443
 biliary stasis in 43
 blood findings in 49
 convalescent phase laboratory
 findings 49
 symptoms 49
 course 440 44
 definition 46
 destruction of parenchyma 49
 430
 preservation of reticulum
 431
- Hepatitis infectious—Cont d
 differentiated from cancer of
 bile ducts 500
 edema of colon 437
 etiology 426
 fibrosis section view 436
 icteric phase 138
 laboratory findings 439
 symptoms 48
 laboratory findings 439
 liver function tests in 439
 lobule destruction view 434
 pathology 464 39
 pre-icteric phase laboratory
 findings 439
 symptoms 438
 preserved reticulum section
 views 432
 prognosis 440
 regeneration of liver cells 433
 of parenchyma section view
 428
 section views 428 43 434 43,
 shrinkage and absorption view
 435
 specific treatment 443
 symptoms 48
 thymol flocculation test 405
 treatment 44
 virus behavior compared with
 homologous serum
 jaundice table 42,
 suppurative 454 477
 complications 46
 diagnosis 476
 etiology 475
 pathology 475
 symptoms 476
 treatment 477
- Hepatocellular jaundice 415
 abnormalities of bile pigment
 metabolism 409
- Hepatoduodenostomy 499
- Hepatolenticular degeneration 474
- Hernia diaphragmatic 115
 types 116
 esophageal hiatus 116
 types illustrated 118
 examination for 37
 inguinal 51
 of small intestine 51
- Herpes labialis allergic aspect 622
- Hexylresorcinol in hookworm 588
 in whipworm 592
- Hiatus hernia esophageal 116
 differentiated from cancer of
 esophagus 111
 insufficiency 116
 pathologic process 117
 varieties 117

- High caloric diet in
 hyperinsulinism 508
 cecum schematic drawing 09
 protein diet for peptic ulcer 171
- Hippuric acid synthesis test of liver
 function 412
- Hirschsprungs disease 310 312
 colectomy subtotal 312
 diagnosis 311
 due to achalasia 312
 treatment 311
- Histamine relation of achlorhydria
 to age shown by 131
 test of gastric function 45
- History 18 7
- Hollander's (insulin) test for gastric
 secretory activity 46
- Homologous serum hepatitis pro
 phylaxis 44? 443
- Hookworm 586 589 *See also*
 Uncinaria
 life history and mode of infection
 587
- Hormones in fatty metamorphosis
 of liver 462
- Horner's syndrome diagnostic
 significance 29
 in carcinoma of esophagus 108
- Hourglass contraction in peptic
 ulcer 210
- Hydatid diseases symptoms of
 Taenia echinococcus 562
- Hydrochloric acid role in peptic
 acid 145
 test of gastric function 48
- Hyperacidity gastric 126
- Hyperalgesia gastric 1 7
- Hyperchlorhydric cases graded
 values 43
- Hyperemia diffuse in bacillary
 dysentery 333
- Hyperesthesia gastric 127
- Hyperfixation of cecocolon 310
- Hyperinsulinism 507
- Hyperplastic stenosing enteritis
 280
- Hyperproteinemia in pyloric
 obstruction treatment
 208
- Hyperprothrombinemia in
 obstructive jaundice 418
- Hyperthyroidism digestive
 symptoms 600
- Hypertrophic biliary cirrhosis 473
 gastritis pathology 137
 symptoms 1-9
- Hyperalbuminemia 403
- Hypochromic microcytic anemia in
 steatorrhea 495
- Hypoglycemia adrenal 523
 epilepsy simulating 5-4
 exclusion in diagnosis of tumor
 of islands of Langerhan
 573
- Hyposensitization in treatment of
 allergies 629
- Hypothyroidism digestive*
 symptoms 600
- Hysterical dysphagia 101
- I
- Icteric phase of infectious hepatitis
 438
- Icterus *See also* Jaundice
 in obstructive jaundice 417
- Ileitis distal and ulcerative colitis
 sketch 342
 regional 283 293
 definition 283
 diagnosis 289 291
 differentiated from acute
 appendicitis 289
 etiology 288
 fistula formation in 289
 giant cell infiltration
 illustrated 287
 granulomatous infiltration
 illustrated 285 286
 ileosigmoidostomy illustrated
 284
 jejunal involvement view 29?
 pathology 283
 prognosis 291
 roentgen view 290
 string sign 291
 symptoms 88
 treatment 292
 ulcerated areas illustrated 283
 stenosing surgery in 93
 terminal 280
- Ileocecal tuberculosis illustrated
 71
- Ileojejunitis medical treatment 20
- Ileosigmoidostomy in regional
 ileitis illustrated 284
- Ileostomy diet following 203
 in ulcerative colitis 361
 postoperative treatment 62
- Ileum diverticula of 248
- Ileum of ulcer *See* Jejunal
 ulcer
- tuberculous ulcers of illustrated
 269

- ileus 6 61
 adynamic 260
 clinical classification and treatment 59
 diagnosis 59
 drugs in treatment 60
 etiology 56
 statistics 2 7
 neurologic 30
 pathology 257
 symptoms 58
 treatment 60
 immunologic aspects of allergies 618
 impaction rectal prevention 20
 incidence of digestive complaints among private patients table 20
 infections oral 818
 infectious hepatitis 426 440 *See also under Hepatitis*
 inferiority *See* Constitutional inferiority
 infusoria 50 54
 ingestion factors interfering with 84
 inguinal hernia 201
 inspection 32
 insulin test of gastric function 45
 intracinar pancreatitis pathology 514
 interlobar pancreatitis pathology 514
 intestinal flukes 580
 lipodystrophy 300
 obstruction 2 6 261
 clinical classification and treatment 59
 diagnosis 59
 etiology 2 6
 statistics 2 7
 incidence 2 6
 pathology 51
 symptoms 258
 treatment 60
 parasites 541 594 *See also* Protozoa Flatworms Round worms
 protein diet 308
 test meals 55
 tuberculosis 268 2 3
 diagnosis 270
 etiology 268
 healing 69
 pathology 268
 symptoms 270
 treatment 70
 intestine amyloid disease of 274
 changes in infectious hepatitis 434
 endometriosis 381
 lardaceous disease of 2 4
 small disorders of pain in 20
 syphilis of 273
 intragastric drips in peptic ulcer 183
 tension measurement 154
 intrahepatic biliary obstruction 413
 intralobular inflammation in infectious hepatitis 427
 intubation in ileus 260
 irritable colon. *See* Colon unstable
 differentiating from gallstones 493
 islands of Langerhans tumors of 5 25 30 *See also* 111
 Tumors
- J
- jaundice 414 422
 bilirubin to study intensity 406
 classification 414 415
 definition 414
 differential diagnosis 413
 hemolytic 415 4 0 4 2
 abnormalities of bile pigment metabolism 409
 definition 4 0
 diagnosis 4 2
 etiology 420
 laboratory findings 4 1
 pathology 420
 symptoms 4 0
 treatment 422
 hepatocellular 415
 abnormalities of bile pigment metabolism 409
 in cancer of pancreas 529
 mechanism diagram 202
 in choledocholithiasis 502
 infectious 444 447
 complications 447
 definition 444
 diagnosis 444
 etiology 445
 geographic distribution 444
 incidence 444
 laboratory findings 44
 pathology 445
 prognosis 447
 stages 446
 symptoms 447
 treatment 447
 intensity bilirubin to study 406
 latent tests for 413

- High caloric diet in
 hyperinsulinism 508
 cecum schematic drawing 309
 protein diet for peptic ulcer 171
- Hippuric acid synthesis test of liver
 function 412
- Hirschsprung's disease 310 31
 colectomy subtotal 312
 diagnosis 311
 due to achalasia 12
 treatment 311
- Histamine relation of achlorhydria
 to age shown by 131
 test of gastric function 45
- History 18 27
- Hollander's (insulin) test for gastric
 secretory activity 46
- Homologous serum hepatitis pro
 phylaxis 442 443
- Hookworm 586 589 *See also*
 Uncinaria
 life history and mode of infection
 587
- Hormones in fatty metamorphosis
 of liver 469
- Horner's syndrome diagnostic
 significance 29
 in carcinoma of esophagus 108
- Hourglass contraction in peptic
 ulcer 219
- Hydatid diseases symptoms of
 Taenia echinococcus 569
- Hydrochloric acid role in peptic
 acid 145
 test of gastric function 48
- Hyperacidity gastric 126
- Hyperalgesia gastric 1-7
- Hyperchlorhydric cases graded
 values 43
- Hyperemia diffuse in bacillary
 dysentery 333
- Hyperesthesia gastric 1 7
- Hyperfixation of cecocolon 10
- Hyperinsulinism 507
- Hyperplastic stenosing enteritis
 280
- Hyperproteinemia in pyloric
 obstruction treatment
 208
- Hyperprothrombinemia in
 obstructive jaundice 418
- Hyperthyroidism digestive
 symptoms 600
- Hypertrophic biliary cirrhosis 413
 gastritis pathology 137
 symptoms 139
- Hypoalbuminemia 403
- Hypochromic microcytic anemia in
 steatorrhea 295
- Hypoglycemia adrenal 593
 epilepsy simulating 524
 exclusion in diagnosis of tumor
 of islands of Langerhan
 593
- Hyposensitization in treatment of
 allergies 629
- Hypothyroidism digestive
 symptoms 600
- Hysterical dysphagia 101
- I
- Icteric phase of infectious hepatitis
 438
- Icterus *See also* Jaundice
 in obstructive jaundice 417
- Ileitis distal and ulcerative colitis
 sketch 342
 regional 283 293
 definition 283
 diagnosis 289 291
 differentiated from acute
 appendicitis 389
 etiology 988
 fistula formation in 89
 giant cell infiltration
 illustrated 87
 granulomatous infiltration
 illustrated 285 86
 ileosigmoidostomy illustrated
 284
 jejunal involvement view 292
 pathology 283
 prognosis 291
 roentgen view 990
 string sign 291
 symptoms 288
 treatment 292
 ulcerated areas illustrated 83
 stenosing surgery in 293
 terminal 80
- Ileocecal tuberculosis illustrated
 271
- Ileojejunitis medical treatment 29
- Ileosigmoidostomy in regional
 ileitis illustrated 84
- Ileostomy diet following 253
 in ulcerative colitis 361
 postoperative treatment 6
- Ileum diverticula of 248
 fistula of 53
 jejunum of ulcer *See* Jejunal
 ulcer
 tuberculous ulcers of illustrated
 263

- flukes 2 5-61
 adynamic 760
 clinical classification and treatment 29
 diagnosis 9
 drugs in treatment 60
 etiology 6
 statistics 34
 neurologic 533
 pathology 45
 symptoms 208
 treatment 60
 Immunologic aspects of allergies 618
 Impaction rectal prevention 40
 Incidence of digestive complaints among private patients table 73
 Infections oral 5152
 Infectious hepatitis 4 6-447 See also under Hepatitis
 Inferiority See Constitutional inferiority
 Infusoria 33 34
 Ingestion factors interfering with 84
 Inguinal hernia 201
 Inspection 32
 Insulin test of gastric function 43
 Interacinar pancreatitis pathology 514
 Interlobar pancreatitis pathology 514
 Intestinal flukes 80
 Hypodystrophy 300
 obstruction 26-61
 clinical classification and treatment 29
 diagnosis 209
 etiology 26
 statistics 5
 incidence 276
 pathology 27
 symptoms 28
 treatment 280
 Parasites 541594 See also Protozoa Flatworms Round worms
 Protein diet 78
 Test meals 3
 Tuberculosis 682 3
 diagnosis 27
 etiology 68
 healing 269
 pathology 68
 symptoms 210
 treatment 212
 Intestine amyloid disease of 274
 changes in infectious hepatitis 434
 endometriosis 381
 lardaceous disease of 74
 small disorders of pain in 20
 syphilis of 213
 Intragastric drips in peptic ulcer 183
 tension measurement 154
 Intrahepatic biliary obstruction 43
 Intralobular inflammation in infections hepatitis 427
 Intubation in ileus 60
 Irritable colon. See Colon unstable differentiating from gallstones 493
 Islands of Langerhans tumors of 3333 See also Islets
 Tumors
 Jaundice 4144
 bilirubin to study intensity 407
 classification 414415
 definition 414
 differential diagnosis 413
 hemolytic 415 420-42
 abnormalities of bile pigment metabolism 409
 definition 40
 diagnosis 42
 etiology 420
 laboratory findings 41
 pathology 40
 symptoms 40
 treatment 422
 hepatocellular 415
 abnormalities of bile pigment metabolism 409
 in cancer of pancreas 59
 mechanism diagram 23
 in choledocholithiasis 302
 infectious 41447
 complications 447
 definition 443
 diagnosis 44
 etiology 443
 geographic distribution 443
 incidence 444
 laboratory findings 44
 pathology 445
 prognosis 44
 stages 446
 symptoms 44
 treatment 447
 intensity bilirubin to study 406
 latent tests for 412

Jaundice—Cont d

- obstructive 414 415 420
 - bile-pigment metabolism in 407
 - complications 418
 - definition 416
 - etiology 416
 - icterus in 417
 - in cancer of pancreas 5 4
 - laboratory findings 418
 - pathology 416
 - symptoms 417
 - treatment 419 420
- postarsphenamine 448
- spirochetal 444 447 *See also*
 - Jaundice infectious
- symptoms general 415
- test of intensity 406
- vitamin K in 404
- Jejunal feedings in peptic ulcer 184
 - ulcer diagnosis 213
 - definition 211
 - etiology 211
 - pathology 211
 - postoperative study 214
 - symptoms 212
 - treatment 215
- Jejunum diverticula of 243
 - fistula of 253
 - primary ulcer 210

K

- Katayama disease 571 577 *See also*
 - Schistosomiasis from *S japonicum*
- Kayser Fleischer corneal ring in
 - hepatolenticular degener-
ation 474
- Ketosis diabetic digestive
 - symptoms 599
- Kidney changes in infectious
 - hepatitis 477
- Kollonnychia in hysterical dysphagia
102

L

- Lambia 552
- Langerhans tumors of islands of
522 525 *See also under*
Tumors
- Lardaceous disease of intestine 274
- Lead colic 601
 - leather bottle stomach 224
- Leg ulcers healing upon improve-
ment of colitis 351
- Leiomyosarcoma of stomach
 - specimen 230
- Leptospirosis icterohaemorrhagica
444 447 *See also*
 - Jaundice infectious

- Leukemia digestive symptoms of
599
- Libman's test 36
- Linitis plastica, 294
- Lipodystrophy intestinal 200
- Lipoid metabolism in liver 403
- Lipophagia granulomatosis 300
- Lipotropic substances 461
- Lithiasis pancreatic 520-527 *See*
also Pancreatic lithiasis
- Liver abscess 415 417
 - adenocarcinoma of from urinary
rectal cancer 372
 - amebiasis involving 547
 - amebic abscess progress 546
 - specimen view 545
- anomalies 401
- bilirubin metabolism in 406
- blood vessels affections 450-455
- cancer 477 481
 - diagnosis 481
 - etiology 477
 - geographic distribution 418
 - table 419
- laboratory findings 480
- pathology 417
- portal cirrhosis and 418
- prognosis 481
- symptoms 479
- treatment 481
- cells necrosis in infectious
 - hepatitis 433
- cholesterol metabolism in 405
- congestion chronic passive 400
- detoxifying function 412
- disease diagnosis direct methods
of 414
- diseases of 401 485
- enlargement in cancer of
pancreas 530
- excretory function 411
- extract crude intravenous in
portal cirrhosis 410
- failure acute course 440
- fat metabolism in 405
- fatty metamorphosis 460 46
- flukes 578 580
- function tests 401 414
 - cephalin cholesterol
flocculation 401
 - galactose tolerance 409
 - Hanger 404
 - hippuric acid synthesis 41
 - in infectious hepatitis 439
 - in portal cirrhosis 469
 - Maclagen 404
 - miscellaneous 412
 - thymol turbidity 404
 - urobilinogen 410
 - use and value 413

- Liver function tests—Cont d
 Van den Bergh 406
 Wallace and Diamond 410
 functions of 401 414
 glucogenesis and glycogenolysis 402
 involvement in amebiasis
 diagnosis 550
 lipid metabolism in 402
 nutmeg 450-452
 phosphatase excretion 411
 protein metabolism in 403 405
 prothrombin manufacture in 404
 reserve power 401
 sarcoma of 481
 shrinking in schistosomiasis
 japonica 575
 tumors 477 482
 benign 477
 Lobular disarray in infectious
 hepatitis 427
 Loop-flotation brine method
 technic 557
 Low cecum 307
 Low fat low cholesterol diet 495
 Lungs examination of 39
 Lymph nodes in neck diagnostic
 significance 0
 Lymphadenitis mesenteric differ-
 entiated from acute
 appendicitis 389
 Lymphopathia venereum anorectal
 95-399
 diagnosis 397 399
 pathology 395 396
 roentgen views 398
 symptoms 97
 treatment 99
 of intestine 274
 Lymphosarcoma diagnosis 65
- M
- Maclagen test 404
 Malignant degeneration in gastric
 ulcer 20* 203
 Marginal jejunal ulcer 211
 Marsupialization of pancreatic cyst
 519
 Mastication disorders of 49
 Maternal overprotection 606
 Mazamorra See Uncinaria
 Meckels diverticulum 249 251
 diagnosis 250
 pathology 449
 symptoms 50
 Megacolon congenital (Hirsch-
 sprung's disease) 311
 Melena in cancer of pancreas 530
 Meniscus sign carcinoma suggested
 by 2 4
 Menopause chronic hepatitis after
 443
 Mental hygiene in constipation 1
 Mercuric chloride poisoning of
 stomach 135
 Merycism 126
 Mesenteric embolism and throm-
 bosis 255
 lymph node tuberculosis 273
 lymphadenitis differentiated from
 acute appendicitis 89
 Metabolic diseases digestive
 symptoms of 599
 Metabolism carbohydrate in liver
 402
 cholesterol in liver 402
 Metagonimus yokogawai 580
 Metastasis of gastric cancer organs
 involved 0
 supraclavicular table of
 incidence 31
 through blood vessels prognostic
 importance 381
 Metastatic cancer of stomach 219
 Meteorism in steatorrhea 294
 Morphine in hemorrhage from
 peptic ulcer 196
 Mother gastrointestinal association
 with 607
 Mother child relationship 606
 Motor test meal 49
 Mouth See also Oral
 diseases of 992
 examination of 29
 lesions in ulcerative colitis 352
 neurotic disorders 90
 Mucin therapy in peptic ulcer 186
 Mucocoele of appendix 391
 Mucous colitis 328 329
 diagnosis 329
 treatment 329
 Mumps oral manifestations 31
 Muscular rigidity abdominal
 tenderness with diag-
 nostic significance 34
- N
- Napl in ring carcinoma of colon
 roentgen view 316
 Nasal septum perforation diag-
 nostic significance 29
 Nausea 127
 diagnostic significance 22
 Necator americanus 586 See also
 Uncinaria
 Neck examination of 30

- Necrosis of liver cells in infectious hepatitis 433
- Needle biopsy for direct study of liver disease 414
- Nematodes 582 594
- Neoplasm hereditary tendency 17
- Neoplastic masses palpation of 33
- Nephritic stage of infectious jaundice 446
- Nervous diseases digestive symptoms in 595 596 factors in constipation 314 tension in ulcerative colitis treatment 615
- Neurologic ileus 595
- Neurosis characteristics of 608 differentiated from psychosis 608 Freud's approach to 605
- Neurotic disorders of mouth 90 patient gastrointestinal symptoms in 608
- Neutral fat absorption in idiopathic steatorrhea 233 red in test of gastric function 47 differential diagnosis by 47
- Nondescent of cecum 508
- Nonrotation of colon schematic drawing 310
- Nose examination of 23
- Nutmeg liver 450 452
- Nutritional deficiencies primary and secondary 84
- Nutritive requirements factors interfering with 85
- O**
- Obstruction intestinal 56 261 *See also* Intestinal obstruction pyloric in peptic ulcer 97
- Obstructive jaundice 414 415 416 420 complications 418 definition 416 etiology 416 hyperprothrombinemia in 418 icterus in 417 laboratory findings 418 pathology 416 symptoms 417 treatment 419 420
- Odor of breath diagnostic significance 29 peculiar to liver disease 467
- Oleoresin aspidii for tapeworm 559
- Omentopexy for portal cirrhosis 471
- Opaque enema 65 66 in unstable colon findings 325 organic disease of colon shown by 65 meal and opaque enema roentgen findings illustrated 63
- Oral *See also* Mouth excretions allergic states and 80 manifestations of deficiency disorders 89 90 sepsis 81 82
- Oriental schistosomiasis 571 577 *See also* Schistosomiasis from *S. japonicum*
- Osteomyelitis of clavicle and ulcerative colitis view 550
- Oxuris vermicularis* 590
- P**
- Pain 50 59 *See also* Abdominal tenderness and pain abdominal diagnostic significance 33 37 characteristics 21 colicky diagnostic significance 36 degree of measurement 56 deep experimental observations 34 effect on position or movement diagnostic significance 2 in acute pancreatitis 510 in cancer of pancreas 598 in colon disorders 20 in duodenal cancer 247 in esophageal lesions 20 in peptic ulcer 150 155 mechanism 152 in perforated peptic ulcer 900 location 20 periodicity diagnostic significance 91 referred 54 severity 21
- Pain food relief cycle in gastric ulcer 150
- Palmer's (hydrochloric acid) test of gastric function 48
- Palpation of abdomen 52 of rectum 37
- Pancreas acute necrosis pathology 510 anomalies 506 azotorrhea in disorders of 507 benign cysts 516 519 classification 516 contents of cavity 518 diagnosis 518 etiology 516 locations diagrams 517

- Pancreas benign cysts—Cont'd
 pathology 516
 prognosis 519
 symptoms 518
 treatment 519
 body of cancer signs and
 symptoms table 5 9
 cancer of 25 39
 ascites 531
 blood signs 531
 creatorrhea as symptom 5 4
 duration 531
 diagnosis 5 6 5 8
 digestive symptoms 5 0
 fecal urobilinogen 534
 feces in characteristics 531
 gallbladder distention 30
 glucose tolerance test 531
 hematemesis in 500
 jaundice as symptom 3 9
 diagram of mechanism 5 2
 laboratory findings 31
 liver enlargement 5 0
 metastasis routes 5 6
 pain 5 8
 pathology 3 6 5 7
 radical operations for sketch
 537
 roentgen examination 534
 serum lipase in 534
 sites of metastasis and invasion
 table 5 5
 symptoms 5 7 31
 treatment 5 8
 weight loss 5 7
 diseases of 06-40
 head of cancer signs and
 symptoms table 5 7
 resection 8
 insufficiency 506
 pseudocyst pathology 518
 resection of head or tail in cancer
 528
 retention cysts pathology 518
 secretions of disorders 306
 Pancreatic duct atypical 506
 edema acute pathology 510
 ferments study in cancer of
 pancreas 534
 function tests for 49 52
 lithiasis 05 2
 diagnosis 520
 roentgen view 521
 symptoms 5 0
 treatment 521
 varieties 5 0
 steatorrhea 298
 tissue accessory of
- Pancreatitis acute 508 313
 diagnosis 31
 differential diagnosis 513
 etiology 508
 Gray Turner sign in 310
 pathology 509
 prognosis 51
 symptoms 510
 treatment 51
 chronic 51 316
 diagnosis 513
 differentiated from cancer of
 pancreas 308
 etiology 513
 hemochromatosis in 515
 pathology 514
 symptoms 514
 treatment 513
 interacinar pathology 14
 interlobar pathology 514
 Iapanicolaou method 24
 Paralysis of esophagus 111
 Parasites intestinal 541 594 8
 also Protozoa Flatworms
 Roundworms
 Parasitic ova diagnosis technic 33
 Parenchymal destruction in infec-
 tious hepatitis 429 4 0
 regeneration section view 4 8
 Parenchymatous disease tests for
 413
 Parorexia 1-8
 Parotitis epidemic oral manifesta-
 tions 91
 Paroxysmal peritonitis benign 6 4
 Pellagra diagnostic signs 30
 Pelvis examination of 8
 Penicillin in ulcerative colitis 64
 Peptic esophagitis primary 100
 ulcer 145 190
 antral gastritis in 146
 atropine derivatives in 146
 bed rest for 170
 belladonna in 146
 benign massive illustrated 149
 of lesser curvature 164 1
 brain lesions in 147
 chronic pathology 148
 complications 191 210
 convalescent diet 174
 deep pain in 36
 diagnosis 134 166
 diets 170 149
 convalescent 144
 high protein 171
 management 134
 principles 140
 Sippy 142
 table of 144

Peptic ulcer—Cont d

- differentiation 166
 - from cancer of stomach 167
 - from colitis 168
 - from dyspepsia 167
 - from gallstones 493
- emotional background of 611
- etiology 145 147
- food for relief of pain 151
- gastrectomy subtotal 180
- gastric contents in 155
- gastritis and 140
- gastroenterostomy in 180
- general directions for patient 175
- healing process 149
 - time 150
- hematemesis 155
- hemorrhage from anatomic lesion how demonstrated 193
 - blood transfusion 195
 - volume in 193
 - diagnosis 192
 - diet 194
 - mortality statistics 199
 - prognosis 197
 - recovery criteria 198
 - symptoms 191
 - treatment 194 199
 - drugs 196
 - surgical 197
- high protein diet 171
- histologic features 149
- hospitalization for 170
- hourglass contraction in 210
- hydrochloric acid in pathogenesis of 145
 - immunity through entero gastrone 185
- incidence 145 147
- intra gastric drips 183
- of esophagus 99 100
- pain in 150 155
 - location 150
 - mechanism 150
 - onset 150
 - relief 151
- pathogenesis 145 147
- pathology 145 150
- perforated 199 201
 - diagnosis 201
 - differentiated from penetrating ulcer 199
 - etiology and incidence 200
 - mortality statistics 201
 - pain 200
 - prognosis 201
 - symptoms 200
- personality types in 169

Peptic ulcer—Cont d

- prognosis 168
- protein hydrolysates 171
- psychogenesis 611
- psychogenic factors 147
- pyloric obstruction in 204 210
 - clinical symptoms 207
 - diagnosis 207
 - differential diagnosis table 205
 - prognosis 208
 - symptoms 204
 - treatment 208
- renal function changes in 206
- sedation 177
- Sippy diet modified 172
- surgical treatment 179 183
 - indications for 180
- symptoms 150 157
- trauma in 147
- treatment 169 186
 - aspiration of stomach 185
 - drugs 175-177
 - enterogastrone 184
 - jejunal feeding 184
 - mucin therapy 186
 - psychosomatic 169
 - radiation therapy 186
 - surgical 179 183
 - types of 169
- urea nitrogen loss 206
- vagus nerve resection 181
- vitamins in 177
- vomiting as symptom 155
- Perforated peptic ulcer 199 201 *See also under* Peptic ulcer
- Periarteritis nodosa 204
- Periportal inflammatory infiltration in infectious hepatitis 4 7
- Periproctitis 393
 - treatment 394
- Peristalsis in scleroderma of esophagus 105
 - reverse syndrome of 103
- Peritonitis benign paroxysmal 694
- Perièche 87
- Pernicious anemia achlorhydria
 - associated with 132
 - digestive symptoms of 599
- Personal history items in 19
- Personality types in peptic ulcer 169
- Phlegmonous gastritis acute 1 5
- Phosphatase excretion by liver 411
- Phthisis dyspepsia of 598
- Phytobezoars 256
- Pica 128
- Piles *See* Hemorrhoids
- Pinworm 290
 - mode of infection 590
 - treatment 591

- Pituitary hypoglycemia 573
 Plumbism diagnosis 601
 symptoms 601
 treatment 602
 Plummer-Vinson syndrome 101
 Pneumatosis gastric 294
 Pneumonia digestive aspects 698
 Poisons in oral excretions 80
 Polycythemia vera digestive
 symptoms of 599
 Polyp benign view of facing 60
 malignant view of facing 60
 Polypoid cancer of colon and rectum
 pathology 370
 of rectosigmoid roentgen view
 377
 of sigmoid colon roentgen
 view 376
 of stomach roentgen views 2-3
 Polyposis of colon hereditary 368
 view of specimen 369
 of rectum simple proctoscopic
 view facing 60
 Polyps gastric 7
 of colon 368
 Pork tapeworm 560
 Portal cirrhosis 467-471 *See also*
 Cirrhosis
 obstruction collateral circulation
 in 466
 diagram 465
 vein diseases of 452-454
 thrombosis of 457
 Postarsphenamine jaundice 448
 Postoperative ulcerative colitis
 treatment 367
 Pregnancy ruptured tubal differ-
 entiated from acute
 appendicitis 389
 Preicteric phase of infectious
 hepatitis 438
 Prepatent stage of schistosomiasis
 578
 Prepyloric ulcer 178
 Proctitis 393
 treatment 394
 Proctoscopic appearance of lesions
 facing 60
 findings normal at various levels
 59
 Proctoscopy 57-60
 in diarrhea 2-9
 position of patient illustrated 58
 technic 59-60
 Proctosigmoidoscopic appearance
 of lesions facing 60
 Proctosigmoidoscopy in cancer of
 rectum 75
 Projectile vomiting significance of
 23
 Prostate disease digestive
 symptoms 601
 Protein diet intestinal 58
 hydrolyzates in peptic ulcer 171
 metabolism in liver 403-405
 Prothrombin manufacture in liver
 404
 Protozoa intestinal 541-555
 Proximal colon fixation of 309
 Pruritus in obstructive jaundice
 treatment 419
 Pseudocyst of pancreas pathology
 518
 Pseudomyxoma peritonei 32
 Psychiatric aspects of digestive
 diseases 603-617
 general concepts 603
 Psychic factors in constipation 314
 in unstable colon 3
 Psychogenesis of peptic ulcer 147
 611
 of ulcerative colitis 612
 Psychosomatic approach to peptic
 ulcer 169
 Psychotherapy in ulcerative colitis
 59-614
 in unstable colon 3-8
 questioning the patient 615
 Psychotic differentiated from
 neurotic 608
 Pteroylglutamic acid in sprue 300
 Pulmonary tuberculosis digestive
 disorders from 32
 Pulsion diverticula of esophagus
 93-94
 Pyelitis acute differentiated from
 acute appendicitis 389
 Pylephlebitis adhesive 452
 suppurative 454
 view 453
 Pyloric end of stomach in peptic
 ulcer pathology 149
 obstruction in peptic ulcer 04
 anatomic lesion determina-
 tion 201
 blood chemistry changes
 206
 clinical symptoms 207
 diagnosis 207
 differential diagnosis table
 205
 hyperproteinemia in 08
 prognosis 308
 symptoms 04
 treatment 08
 vomiting an indicator of 3
 Stenosis congenital 124
 treatment 1-5

Q

Quincke's disease 63

R

Radiation therapy in peptic ulcer 186
 Raynaud's disease signs in scleroderma of esophagus 105
 Rectal constipation 316
 varieties 317
 mucosa examination for schistosoma japonicum 516
 pain 21
 Rectum adenocarcinoma specimen 371
 cancer of 369 381 *See also under*
 Cancer
 diseases of 93 400
 examination of 37
 fissure of 394
 fistula of 394
 impaction of prevention 320
 Red cell fragility in hemolytic jaundice 421
 Redundant colon 304 307 *See also under* Colon
 Referred pain 34
 Regeneration of liver cells in infectious hepatitis 433
 Regional ileitis 283 293
 Regurgitation 24
 Renal colic differentiation from
 gallstones 493
 disease organic digestive symptoms 600
 function changes in peptic ulcer 406
 Resection gastric achlorhydria associated with 132
 for cancer of stomach 299
 of head of pancreas 538
 Respiratory diseases digestive aspects of 597
 Retention cysts of pancreas pathology 518
 Retroperitoneal hernia 251
 Reverse peristalsis syndrome of 23
 Riboflavin deficiency diagnostic signs 30
 glossal clinical picture 87 90
 illustrated 83
 Riedel's lobe 401
 Roentgen examination 60 67
 routine normal findings chart 62
 Rose bengal test 412

Roundworms 581 594 *See also*
Ascaris lumbricoides
 Rumination 126

S

Salivary calculi 91
 system disorders of 90 92
 Salpingitis differentiated from acute appendicitis 389
 Santonin for *Ascaris lumbricoides* 590
 Sarcoma of liver 481
 of stomach 229 232
 diagnosis 231
 esophagoscopy in 231
 incidence 229
 pathology 229
 symptoms 231
 treatment 32
 Schatzki's small bowel enema in tumors of small intestine 465
 Schistosoma common features 564
 general description 564 566
 life cycle diagram 565
 types 564
 Schistosomiasis from *S. haematobium* 577 518
 from *S. japonicum* 515 577
 acute stage 574
 asymptomatic infection 574
 chronic stage 515
 complications of acute phase 574
 diagnosis 576
 etiology 571
 fulminating infections 514
 geographic distribution 571
 incubation stage 513
 insidious infection 574
 mode of infection 571
 pathology 571
 section view of liver 519
 symptoms 573
 treatment 577
 systemic manifestations 573
 from *S. mansoni* 566 571
 cirrhosis of liver view 467
 description 566
 diagnosis 510
 geographic distribution 566
 mode of infection 566
 pathology 566
 prophylaxis 510
 stages 568 569
 symptoms 568
 treatment 570
 oriental *See* Schistosomiasis
 from *S. japonicum*

- Schmidt intestinal test diet 55
- Scleroderma of esophagus 104 107
 diagnosis 105
 etiology and pathology 104
 of lesions illustrated 106 107
 symptoms 105
 treatment 105
 types 105
- Scorbutic gums illustrated 83
- Secretin test of pancreatic function
 49
 differential signs 51
 reaction types 50
 technic 50
- Secretory test Hollander's insulin
 46
 meals 46
- Security threat to intensification
 of gastrointestinal
 symptoms 610
- Sedation in gastric irritation 1 0
 in peptic ulcer 177
- Sensation gastric disorders of 1 7
- Septicemic stage of infectious
 jaundice 446
- Serum alkaline phosphatase test of
 liver function 411
 amylase concentration in acute
 pancreatitis 512
 determination in perforated
 peptic ulcer 401
 bilirubin in hemolytic jaundice
 421
 lipase in cancer of pancreas 534
- Sialadenitis chronic 91
- Sialodochitis 91
- Sialolithiasis 91
- Sialorrhea 90
- Sickle cell anemia digestive
 symptoms of 599
- Sigmoid colon polypoid carcinoma
 roentgen view 476
- Sippy diet 170
 modified 172
- Skin examination of 27 49
 signs of Addison's disease 28
 tests for gastrointestinal allergies
 6 7
- Small intestine atresia of 49
 diseases of 241 303
 fistulas of 52 256
 hernia of 251
 tumors of 267
 diagnosis 265
 pathology 263
 treatment 66
 volvulus of 261
 diagnosis 26
- Sore tongue 90
- Spasm of esophagus *See under*
 Esophagus
- Spasmophilia 3
- Spastic colon *See also* Colon
 unstable
 constipation in 316
- Spastic phenomena in stomach
 disorders 126
- Spleen changes in infectious
 hepatitis 437
- Splenomegaly in hemolytic jaundice
 421
 in hysterical dysphagia 107
- Sporocyst formation in
 schistosomes 565
- Spreading cancer of stomach 419
- Sprue *See also* Steatorrhea
 idiopathic
 diagnostic signs 30
 folic acid in 400
 pernicious anemia associated
 with 297
 roentgen views 297
 tropical and pernicious anemia
 differentiation with
 neutral red 47
- Stasis biliary in infectious
 hepatitis 433
- Steatorrhea diagnosis 279
 idiopathic 293 300
 calcium deficiency in effects
 295
 definition 49
 diagnosis 298
 diet in 294
 differential diagnosis 98
 etiology 293
 hypochromic microcytic anemia
 in 95
 incidence 293
 meteorism in 294
 mucosal changes 296
 pathology 294
 roentgen findings 96
 segmentation in 298
 stool in 94
 symptoms 294 96
 treatment 299
 vitamin deficiency in 49
 in pancreatic disorders 507
- Stenosis pyloric congenital 1 4
 in adults 125
 treatment 125
- Sterlin's sign in intestinal
 tuberculosis 477

- Stomach** *See also* Gastric
 acute dilatation of 141
 anomalies of 114 125
 aspiration of 42
 in peptic ulcer 185
 barium meal study of 61
 benign tumors of 227
 bubble chronic 24
 cancer of 217 240
 achlorhydria associated with 132
 atrophic gastritis and 217
 diagnosis 222
 differential diagnosis 225
 dyspepsia as symptoms 221
 fungating 218
 gastric analysis in 274
 incidence 217
 table 218
 organs involved in spreading 270
 pathogenesis 217
 pathology 218 220
 polypoid roentgen views 273 226
 prognosis 227
 roentgenologic diagnosis 224
 schematic drawings of various types 222
 specimen 228
 spreading 219
 symptoms 220
 treatment 229
 ulcerated 219
 diseases of 114 240
 diverticulum of 114
 fasting aspiration of 46
 foreign bodies in 236
 hair balls in 236
 hemorrhagic lesions 147
 leiomyosarcoma specimen 230
 mercuric chloride poisoning of 135
 sarcoma of 279 237
 diagnosis 231
 incidence 229
 pathology 279
 symptoms 231
 treatment 237
 syphilis of 232 233
 tube illustration 41
 technic of using 41
 tuberculosis of 233 234
 volvulus of 234
 treatment 236
Stomatitis aphthous diagnosis 80
 treatment 81
Stone *common form in*
 cholelithiasis 489
Stool *See also* Feces
 abnormal 56
 appearance in idiosyncratic steatorrhea 294
 examination in amebiasis 548
 in diarrhea 278
 in obstructive jaundice 418
 in *Schistosoma japonicum* 516
 in spastic constipation 316
 normal 56
 training effect on personality development 607
Stricture of bile ducts 497 499
 of esophagus 101
String sign in regional ileitis 491
Strongyloides stercoralis 592
Subacidity gastric 127
Sulfathaladine in regional ileitis 92
Sulfonamides in ulcerative colitis 360
Suppurative hepatitis 475 477 *See also under* Hepatitis
 pyelephlebitis 474
 view 453
Supraclavicular metastases table of incidence 31
Sympathicotonia 327
Symptoms cardinal 20
 evaluation scheme 27
Syphilis cerebrospinal digestive lesions from 595
 of intestine 273
 of stomach 227 232 233
- T**
- Tabes mesenterica** 213
Tabetic gastric crisis Argyll Robertson pupil in 29
Taenia echinococcus 561 563
 description 561
 diagnosis 562
 prophylaxis 63
 symptoms 567
 treatment 563
Taenia saginata 556 560
 description 556
 diagnosis 557
 prophylaxis 557
 symptoms 557
 treatment 559
Taenia solium 560
Taeniae differential diagnosis table 557
Takata Ara test 412
Talma Morison operation for portal cirrhosis 471
Tapeworm diagnosis 557
 life history and mode of infection diagram 548

- Tapeworm—Cont d
 prophylaxis 557
 symptoms 557
 treatment, 559
- Teeth examination of 29
- Tenderness abdominal diagnostic significance 33 37
 with muscular rigidity diagnostic significance 34
- Test meals 4° 52
 intestinal 55
 motor 49
 odor 44
 procedures not needed 44 45
 secretory 42
- Tetrachlorethylene for whipworm 92
- Threadworm 590
- Thrombosis mesenteric 255
 of portal vein 452
- Thymol in hookworm disease 588
 turbidity test 404
- Tone of abdominal wall 3
- Tongue coated diagnostic significance 25
 examination of 30
 signs in idiopathic steatorrhea 94
- Topfer's reagent in test for gastric acidity 43
- Traction diverticula of esophagus 95
- Transfusion in hemorrhage from peptic ulcer 195
- Trauma in peptic ulcer 147
- Trematodes 564 581
- Trichinella spiralis* description 582
 mode of infection 58
- Trichiniasis diagnosis 583
 incidence 582
 prophylaxis 583
 slide views 584 585
 symptoms 582
 treatment 586
- Trichobezoars 256
- Trichocephalus trichiurus* 59
- Trichomonas intestinalis* 553
- Trichuris trichiura* 592
- Trophozoite noninfectious nature 544
- Tuberculosis abdominal 268 273
 digestive disorders associated with 598
 ileocecal illustrated 71
 of appendix 268
 of colon diagnosis 272
 etiology 68
 healing 269
 illustrated 269
 pathology 268
- Tuberculosis of colon—Cont d
 symptoms 270
 treatment 72
 of mesenteric lymph nodes 13
 of stomach 23 234
 pulmonary digestive disorders from 3
- Tuberculous ulcers of ileum illustrated 269
- Tumor mass in regional ileitis 288
- Tumors of appendix 91
 of colon 67 381 See also Cancer of colon
 benign 367
 of islands of Langerhans 5° 5 5
 diagnosis 5 3
 pathology 522
 symptoms 5°°
 treatment 524
 of liver 477 481
 benign 477
 of small intestine 261 265
 diagnosis 265
 pathology 63
 treatment 265
- U
- Ulcer anastomotic jejunal 211
 anastomotic (marginal) roentgen view 14
 crater 160
 illustrated 158 159 161
 duodenal roentgen views 209
 gastrojejunal 211
 jejunal 211 215
 definition 211
 diagnosis 13
 etiology 211
 pathology 211
 postoperative study 214
 primary 210
 symptoms 212
 treatment 215
 of rectum tuberculous proctoscopic view facing 60
 of second portion of duodenum 219
 peptic See Peptic ulcer
 symptoms emotional background of 611
- Ulcerated cancer of stomach 219
- Ulcerative colitis 43 6 See also Colitis ulcerative
 allergic aspect 6°4
 psychiatric 614
 psychogenesis 613
 psychotherapy 614
- Umbilical hernia 251

- Uncinaria 586 589
 - description 586
 - diagnosis 588
 - prophylaxis 588
 - stages of intestinal parasitism 588
 - symptoms of invasion 586
 - treatment 588
- Unconscious theory of 605
- Unstable colon 32° 328 *See also*
under Colon
- Urea nitrogen retention in peptic ulcer 206
- Ureteral calculus differentiated from acute appendicitis 389
- Urinary urobilinogen excretion 408
- Urobilinogen excretion normal steps leading to 407
- fecal in cancer of pancreas 534
- in hemolytic jaundice 428
- oxidation of 408
- test 410
- Urologic disorders digestive symptoms 600
- Utilization of intake factors interfering with 85

V

- Vagotomy in peptic ulcer complications 182
 - insulin test after 45
 - plus gastroenterostomy 182
 - ulcer pain after relief 155
- Vagotonia 3°
- Vagus nerves resection in peptic ulcer 181
- Van den Bergh reaction 406
- Varices esophageal 102 104
 - diagnosis 104
 - pathologic process 10° 103
 - treatment 104
- Varicose veins at lower end of esophagus view 464
- Veins hepatic diseases of 454
- Vincent's infection 61 62
- Virchow's node diagnostic significance 31
- Visceroptosis 74 76
 - diagnosis 74
 - operations 76
 - treatment 75
- Vitamin B complex deficiency oral lesions 86
- Vitamin C deficiency oral lesions 86
 - scorbutic gums from 83
- Vitamin deficiency in steatorrhea 95
- Vitamin K in prothrombin manufacture 404
- Vitamin requirements in obstructive jaundice 419
- Vitamins in peptic ulcer 177
- Volvulus of small intestine 461
 - diagnosis 26°
 - of stomach 34
 - treatment 236
- Vomiting diagnostic significance °
 - etiology 22
 - in peptic ulcer 155
 - mechanism 22
 - projectile significance 3

W

- Wall infection of intestine 481
- Wallace and Diamond test for urinary urobilinogen 410
- Weight loss in cancer of pancreas 527
 - record importance in examination 48
- Wells disease 444 444 *See also*
Jaundice infectious
- Whipple's disease 300
- Whipworm 592
- Wilson's disease 474
- Wirsung duct of atypical 506

X

- Xanthomas diagnostic significance 28
- Xanthomatous biliary cirrhosis skin signs 28
- Xelostomia 30

